#### DRAFT

REMEDIAL INVESTIGATION REPORT VOLUME 1 OF 2

ECC SITE

ZIONSVILLE SITE

WA18.SL30.0

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#### Chapter 1

#### **EXECUTIVE SUMMARY**

## BACKGROUND

The Environmental Conservation and Chemical Corporation (ECC) site is in Boone County, 865 south U.S. 421, Zionsville, Indiana, about 10 miles northwest of Indianapolis. The site occupies 6.5 acres alongside the 168 acre Northside Sanitary Landfill (NSL), an ongoing solid waste disposal facility. The ECC site is bounded on the south and east by the landfill. An unnamed ditch separates the two facilities along the east boundary. The site is bounded on the north and west sides by several residential homes, located within one-half mile of the facility.

ECC began operations in 1977 and was engaged in the recovery/
reclamation/brokering of primary solvents, oils and other
wastes received from industrial clients. Waste products
were received in drums and bulk tankers and prepared for
subsequent reclamation or disposal. Reclamation processes
included distillation, evaporation and fractionation to reclaim solvents and oil.

Accumulation of contaminated stormwater onsite, poor management of the drum inventory and several spill incidents caused initial state and EPA investigations that later led to civil suits and finally placement of ECC into receivership in July 1981. Drum shipments to the site were halted in February 1982. The company was found insolvent in August 1982 and the state and EPA began plans for cleanup. Numerous site investigations, including sampling and analysis were conducted during the period.

Removal measures at the site began in March 1983 and continued through 1984. Actions included removal and treatment or disposal of cooling pond waters, approximately 30,000 drums of waste, 220,000 gallons of hazardous waste from tanks and 5,650 yd³ of contaminated soil and cooling pond sludge. A clay cover, placed over the site, was recently compacted.

## REMEDIAL INVESTIGATIONS

#### SCOPE

Remedial investigations began in 1983 and continued until December 1984. Soil, hydrogeologic, and surface water and sediment investigations were conducted.

Two phases of soil sampling were conducted. Phase 1 consisted of 15 surficial soil samples and 15 shallow (2.5 foot depth) borings and was conducted before removal of 2 feet of

contaminated surface soil from most of the site. Phase 2, conducted after soil removal, consisted of 9 soil borings (up to 12 feet in depth) through the concrete pad on the south 1/3 of the site and 12 test pits to depths up to 10 feet in the remaining areas.

Hydrogeologic investigations included an electrical resistivity survey, test drilling, monitoring well installation, monitoring well sampling and residential well sampling. A total of 16 2-inch diameter PVC monitoring wells were installed in 3 phases. Wells were placed to monitor the shallow saturated zone, the shallow sand and gravel aquifer and the deep confined aquifer. Groundwater sampling was also performed in 3 phases. In addition, 5 residential wells were also sampled.

Surface water investigations included three onsite and four offsite surface water samples and 6 offsite sediment samples.

## RESULTS

Onsite soil sample inorganic analysis results showed only antimony, cadmium, cobalt, copper, lead, manganese, and zinc were at concentrations exceeding the typical range in soil.

Of these, cadmium, lead, and zinc were reported in more than one sample at concentrations exceeding the typical range in soils. Exceedance of the typical ranges in soil samples of

inorganic constituents beneath the concrete pad is relatively minor relative to the soil contamination in the northern drum and tank storage areas. Inorganic contamination of the soil is apparently greatest in the near surface (0-3 feet) soil in northern portions of the site. Inorganic contamination does appear to extend to depths of at least 5 feet in the northern portions of the site, although it is less widespread than observed in the overlying shallow soil.

Primary organic contaminants found in site soils are volatile organic compounds and phthalates. These compound groups are the most widespread organic contaminants and are generally present in the highest concentrations. Total volatile organic contaminants (VOC's) ranged from 16 to 14,604,000 ug/kg. Total phthalates ranged from "not detected" to 370,000 ug/kg. Organic contamination decreases in the variety of compounds and their associated concentrations with depth. However, organic contaminants were detected to the maximum depth of sample analysis (8.5 feet).

Results of the hydrogeologic investigations indicate the existence of 4 hydrogeologic units in the area, a shallow saturated zone, a shallow sand and gravel aquifer, a silty clay and clayey silt zone and a deep confined aquifer.

Migration of soil contaminants to the shallow saturated zone has occurred onsite as evidenced by high levels of organic contaminants in one well onsite. The shallow sand and gravel aquifer has been shown to be contaminated with inorganics and organics in one well offsite and lesser amounts of organics in one well onsite and another immediately adjacent and downgradient of the site. Because of the presence of the NSL east of ECC, it cannot be definitively stated that the source of offsite contamination is ECC though the contaminants are consistent with those found onsite. Organic contamination in the other two wells is likely due to onsite soils at ECC since they are directly downgradient of ECC contaminated soils and not NSL.

Contamination of the shallow sand and gravel aquifer may have occurred either via migration through the silty clay till onsite or through contaminated water and sediment in the former cooling water pond, since it intersected the shallow sand and gravel aquifer before to its removal and backfilling.

The deep confined aquifer below the site has not been found to be contaminated. Future migration of onsite contaminants to the deep aquifer is highly unlikely due to an upward vertical hydraulic gradient.

Migration of contaminants to the nearest residential wells surrounding the site is not indicated by the results of the residential well sampling.

Surface water sampling results indicate that inorganic contamination of surface water does not appear to be occurring offsite in the vicinity of ECC. Inorganic sediment contamination in the vicinity of ECC is limited to lead in the unnamed ditch. Organic contamination of offsite surface water was found in Finley Creek near Highway 421. Contaminants consist almost entirely of chlorinated hydrocarbons and are likely from ECC. Also, surface water ponded on the clay cap onsite was found to be contaminated with a variety of base/neutrals and volatile compounds.

Two organic compounds possibly resulting from the ECC site were found in sediments in the unnamed ditch and and in Finley Creek near Highway 421.

#### CONTAMINANT TRANSPORT AND FATE

Analytical results of the remedial investigations characterize current site contamination. Future conditions assuming no action is taken at the site were estimated based on potential transport pathways and the natural attenuation and degradation of contaminants. Due to the large numbers of site contaminants, 14 indicator chemicals from four major contaminant groups were used in the estimation of transport and GLT424/52

fate. Transport and fate are briefly summarized here for volatile organic contaminants, phenols, phthalates, and polychlorinated biphenyl's (PCB's). Transport of inorganic constituents from the soil is considered negligible due to the low levels found and the adsorptive capacity of the onsite soils.

All indicator volatiles in the subsurface soils except

1,1,1-trichloroethane, trichloroethene, and tetrachloroethene
are estimated to naturally degrade to acceptable levels (as
calculated in the endangerment assessment) within 10 years.
These three are estimated to take from 10 to 100 years at
average site concentrations and from 100 to 1,000 years at
maximum site concentrations to degrade. Phenols and
phthalates in the subsurface soil are already below
acceptable levels and are estimated to degrade to trace
levels within 10 years and 200 years, respectively. PCB's
in the soil are estimated to take 50 years at average
concentrations and 100 to 2,000 years at maximum
concentrations to degrade to acceptable levels.

Under existing site conditions, the volatiles, phenols, and certain phthalates will tend to leach from subsurface soil into the groundwater and slowly migrate to the unnamed ditch or Finley Creek (PCB's and most phthalates will only leach in trace amounts). Estimates for travel time vary from 20 years to 8,400 years depending upon the compound,

hydraulic conductivity, and travel distance. Once in the surface waters, contaminants will either volatilize, adsorb to sediments, or experience large dilutions before reaching the Eagle Creek Reservoir.

#### ENDANGERMENT ASSESSMENT

The endangerment assessment found that under the no action alternative potential risk to human health and the environment exist at the ECC site. The affected media are soil, groundwater and surface water. They were assessed based on comparison of concentrations at exposure points to lifetime excess cancer risks, acceptable daily intake values, and relevant or applicable standards, criteria or guidelines. For the public health concerns residential and occupational use settings were used in assessing risk. An excess lifetime cancer risk of  $1 \times 10^{-6}$  is often used to reflect a level of concern for carcinogen risk.

For public health concerns, the exposure routes that resulted in an excess lifetime cancer risk greater than  $1 \times 10^{-6}$  are listed below:

o Soil via ingestion: the south concrete pad soil at intermediate depth in a residential setting; and north test pit area at shallow and intermediate depth in residential and occupational use settings.

- o Groundwater via ingestion: the shallow saturated zone and shallow sand and gravel aquifer at current concentrations in both use settings; the shallow saturated zone at future projected concentrations in both use settings.
- o Groundwater via dermal absorption of volatile organic compounds: during bathing, the shallow saturated zone and shallow sand and gravel aquifer at current conditions in the residential setting; the shallow saturated zone at future projected concentrations in the residential setting.
- o Ingestion of fish that bioconcentrated contaminants from the surface water: Finley Creek under the lowest dilution situation at projected concentrations.

Risk from dermal absorption of volatile compounds via wading in the surface water does not exceed  $1 \times 10^{-6}$ . However, wading in the unnamed ditch and in Finley Creek under the lowest dilution situation has excess lifetime cancer risks between  $1 \times 10^{-6}$  and  $1 \times 10^{-7}$ . Given the uncertainty in both risk estimation and fate, and transport calculations, it is possible for the risk to be oders-of-magnitude higher or lower than estimated.

For environmental concerns the projected release of contaminants to the surface water in the unnamed ditch should not exceed the ambient water quality criteria for protection of aquatic life and other known  $LC_{50}$  values.

The risk analysis performed for the endangerment assessment is conservative and tends to reflect upper bound exposures.

However, given the uncertainty in both risk estimation and fate and transport calcuations, the actual risks may be lower or higher than estimated.

The current impact of the site is limited due to the low population at risk. Site location and environmental media characteristics (for example, low groundwater flow velocity) limit the population at risk if there is future development of the site and the surrounding area under the no action alternative. The environmental impacts also would be similarly restricted.

In conclusion, the ECC site does pose a threat to the public health, welfare, and environment and a feasibility study of remedial actions to cost-effectively mitigate the site hazards should be performed.

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### Chapter 2

#### INTRODUCTION

This remedial investigation (RI) report for the Environmental Chemical and Conservation Corporation (ECC) site near Zions-ville, Indiana, is prepared in partial satisfaction of Contract No. 68-01-6692, Work Assignment No. 18.5L30.0, and the Final Work Plan (April 1983), Tasks 1 through 5.

### PURPOSE OF THE REPORT

This RI report is based, in part, on data obtained during remedial investigation activities conducted from April 1983 through December 1984 at the ECC site. These data and those from other sources are used to define the site problems, identify pathways and receptors, and determine the necessity for and extent of remedial actions at the site.

The purpose of this RI report is threefold: 1) document the details of remedial investigation activities through technical memorandums included in Appendix A, 2) summarize and present the site investigation analyses and conclusions, and 3) determine if there is a threat to public health, welfare or the environment.

## ORGANIZATION OF THE REPORT

This RI report is organized into four main sections. Chapter 3 presents a description of the site and its history. Chapter 4 presents the summary and results of the RI. Chapter 5 presents contaminant transport and fate. Chapter 6 presents the methodology and results of the endangerment assessment. Volume 2 of the RI Report presents the appendixes that contain detailed documentation of activities and specific data obtained for each task completed during the RI.

### RI ACTIVITY TECHNICAL MEMORANDUMS

Each RI activity is described in a technical memorandum (TM) issued during the course of RI work. These TM's are contained in Appendix A of this report. Each TM describes specific procedures, observations, measurements, and data results of RI activities.

#### ANALYSIS OF SITE INVESTIGATIONS

The results of site investigations conducted at ECC from April 1983 through December 1984 are organized by the operable units. The analysis provides the technical basis for identification of problems and pathways of contamination for each operable unit.

## CONTAMINANT TRANSPORT AND FATE

The pathways of contamination are identified and estimated ranges of transport rates and fates of contaminants are presented. The results form the basis of the assessment of the no action alternative.

## ENDANGERMENT ASSESSMENT

The results of the site investigations and the contaminant transport and fate analysis are used in the endangerment assessment to determine if a threat to human health or the environment exists at the site. The endangerment assessment will in turn be used in deciding if a feasibility study is necessary at the site and, if so, what the remedial action objectives will be.

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## Chapter 3

#### SITE BACKGROUND

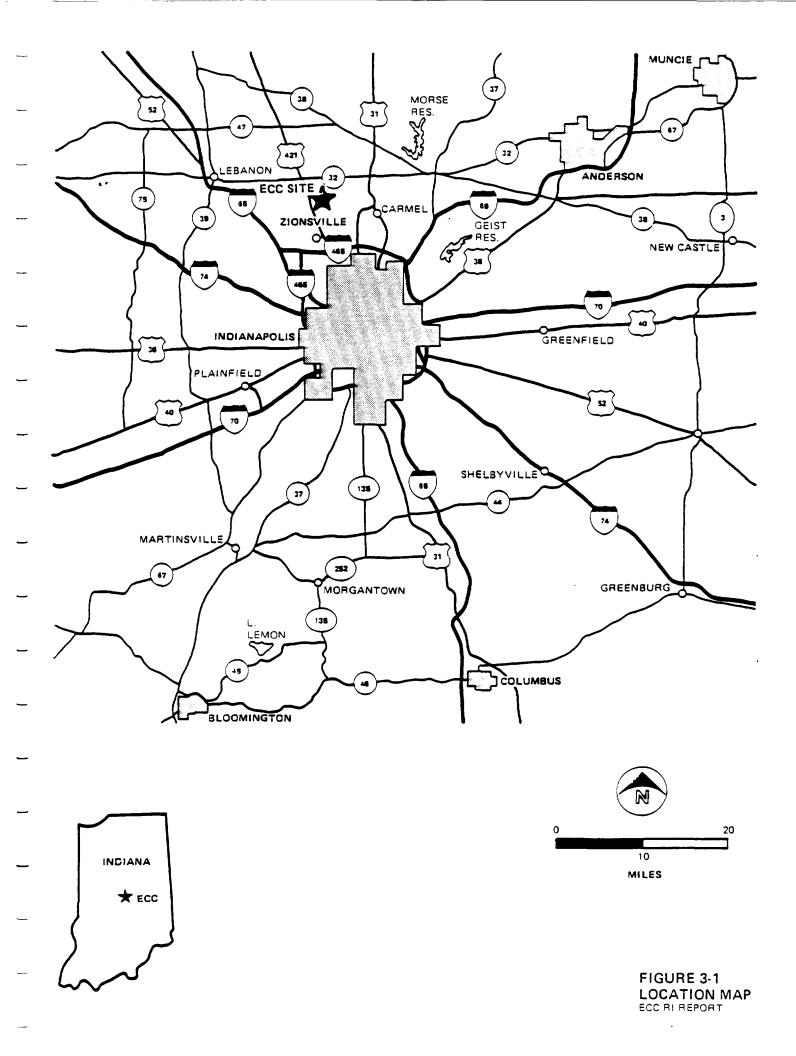
### SITE DESCRIPTION

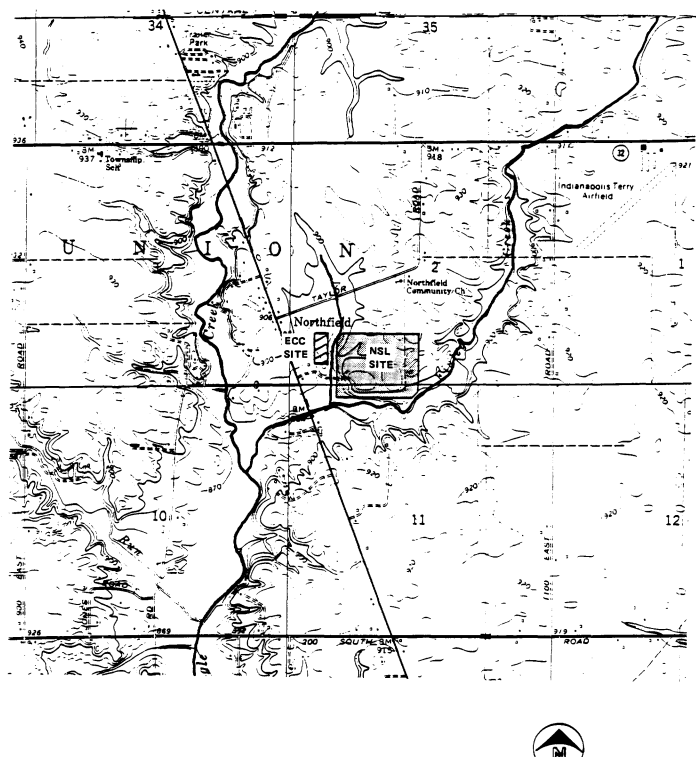
ECC is in Boone County, 865 south U.S. 421, Zionsville, Indiana, about 10 miles northwest of Indianapolis (Figure 3-1). The site occupies 6.5 acres alongside the 168 acre Northside Sanitary Landfill (NSL), an ongoing solid waste disposal facility (Figure 3-2).

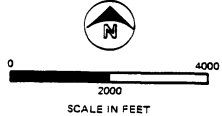
The ECC facility is bounded on the east by the landfill. A site map showing the site as it was in 1982 is shown in Figure 3-3. An unnamed ditch separates the two facilities along the east boundary. The site is bounded on the north and west sides by several residential homes, all located within one-half mile of the facility.

#### SITE HISTORY

ECC began operation in August of 1977 under a construction permit issued by the Indiana Air Pollution Control Department (APCD) on May 5, 1977. The company was engaged in the recovery/reclamation/brokering of primary solvents, oils and other wastes received from industrial clients. Waste products were received in drums and bulk tankers and prepared







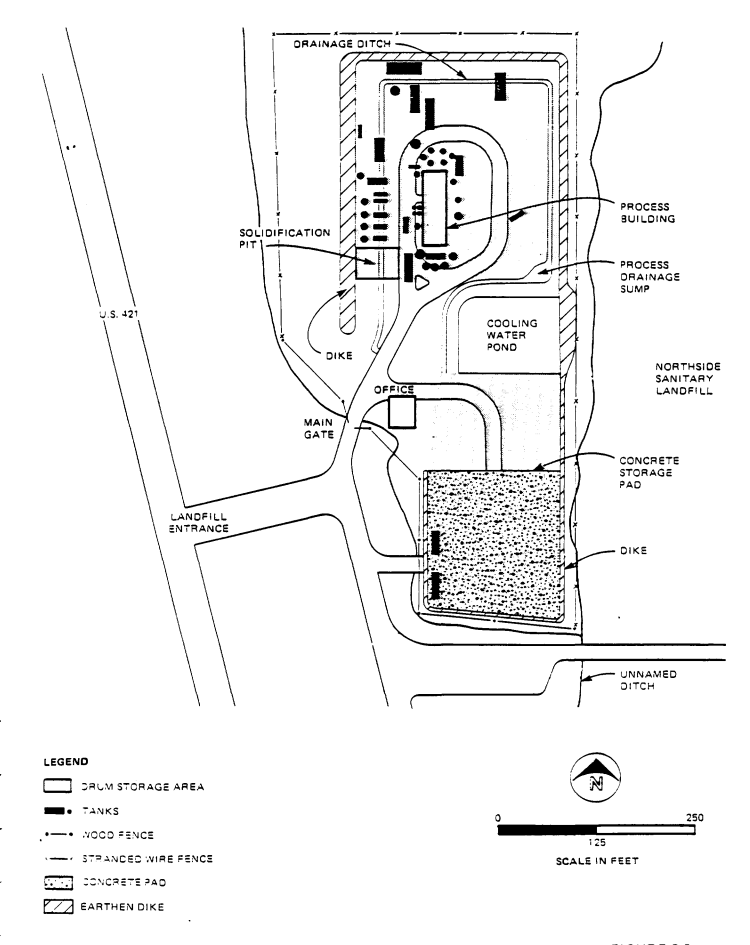


FIGURE 3-3 SITE MAP (1982) ECC RI REPORT

for subsequent reclamation or disposal. Reclamation processes included distillation, evaporation and fractionation to reclaim solvents and oil.

Two problems developed during the facility's operation:

- o The inability of the company to adequately dispose of wastewater and contaminated stormwater generated at the facility,
- o The inability of the company to manage its drum inventory in a manner that did not pose a threat to the environment.

In an attempt to handle the wastes generated onsite, approval was sought by ECC to dispose of 5,000 gallons per day of oil recovery wastes and 1,000 to 1,500 gallons per week of still bottoms at NSL. Approval to dispose of the still bottoms was granted (with conditions) by the SPCB on October 11, 1977; however, the request to dispose of the liquid waste from the oil recovery operations was denied.

Subsequently, the company sought other avenues of waste disposal. An agreement was reached between the Indiana State Board of Health (ISBH), ECC, and NSL to allow disposal of oily wastes in the landfill with municipal refuse. Following expiration of this agreement in May 1979, ECC added units to

process wastewater by distillation onsite. The product water was used as boiler makeup water.

On July 31, 1979, the ISBH received a report from a private citizen that an oil spill had occurred on Eagle Creek north of Zionsville. Immediate inspection revealed that the oil had originated from ECC and a minor amount from NSL. ECC agreed to take action to recover the oil. A followup investigation conducted on August 2, 1979 by the ISBH showed that ECC intentionally discharged process and cooling water from a storage lagoon into Finley Creek without a permit. ECC officials explained that due to heavy rains, stormwater pumped from the drum storage and loading areas to the cooling water pond caused it to overflow. Therefore, it became necessary to drain the excess water.

On September 18, 1979, the SPCB met to discuss the spill and discharge incidents at ECC. The board ratified an Agreed Order that included a fine and provisions to upgrade the methods of recordkeeping at the facility. In November 1979, the SPCB began a water sampling and analysis program at the site. Cooling water pond samples taken on November 2, 1979 were found to contain high concentrations of arsenic, cadmium, chromium, lead, nickel, oil and grease, phenol, and zinc. Further testing of area wells and streams were inconclusive in documenting contamination of groundwater and surface water.

In December 1979, the U.S. EPA designated ECC as a potential hazardous waste site and began investigations under the Hazardous Materials Emergency Response Program. By April 17, 1980, the ISBH submitted documentation to the Indiana Environmental Management Board (EMB) concerning ECC violations of the Environmental Management Act, the Air Pollution Control Law and the Stream Pollution Control Law. Specifically, the staff documented that:

- o ECC posed a threat to pollute the environment.
- o The company was burning chlorinated hydrocarbons and other solvents as boiler fuel without approval.
- o Process water and contaminated stormwater were discharged without approval.
- o Spills of oil and other objectionable substances occurred and were not reported or effectively cleaned up.

Based on these violations, the EMB referred the matter to the Office of the Attorney General on May 15, 1980 for appropriate enforcement.

On February 9, 1981, an ECC employee died of exposure to toxic vapors after entering a solvent tanker.

A Consent Decree was issued on July 1, 1981, by the Boone County Circuit Court imposing a \$50,000 civil penalty against ECC. Furthermore, the court placed ECC into receivership and prohibited the company from using NSL for disposal of wastes. The decree gave ECC until November 1, 1982 to comply with environmental laws and regulations.

At this point, the ISBH began weekly monitoring of ECC's drum storage area to insure that action was being taken to reduce barrel inventory and improve storage facilities. The area was found to be extremely overcrowded with drums, some of which were damaged and leaking. Access was also dangerously poor. By October of 1981, construction of a concrete drum storage pad was underway and drum inventory had been reduced to an estimated 20,000 barrels. By December, the number of leaking, formerly leaking, popped top, corroded/damaged, and bungless/open top drums had been reduced to about 225. In February 1982, the EMB placed a freeze on drum shipments to the facility before the Boone County Circuit Court to assure compliance with the Consent Decree regarding storage of drums, location of materials onsite and in transit, and the removal of sludge.

On May 5, 1982, ECC was ordered by the court to close and environmentally secure the site for failure to reduce hazardous waste inventories. Two days later ECC's court receiver filed a closure plan with the Boone County Circuit

Court. By August 1982, ECC was found to be insolvent and planning work had begun for environmental revitalization, cleanup, and recycling of the site.

On September 21, 1982, the Office of the Attorney General held a conference with the ISBH and representatives from 60 generators of waste to propose a voluntary cleanup plan for the ECC site. The closure plan and settlement offer required generators to remove and dispose of wastes and pay \$250/drum into a trust fund to be used for remaining surface/subsurface remedial measures. In return, generators would receive a limited release. In response to the offer, the generators entered into a loose coalition and hired Chemical Waste Management, Inc., to prepare a technical proposal for a complete surface cleanup. Initial negotiations between U.S. EPA and the generators for site surface cleanup were not successful.

#### PREVIOUS INVESTIGATIONS

Sampling and testing efforts were conducted at ECC from 1976 through 1982. Sources of data were primarily laboratory data sheets or handwritten data summary tables, generally unaccompanied by descriptions of the sampling and testing procedures used. As such, much of this historical data summarized herein could not be used as a basis for definitive interpretations of existing conditions onsite or offsite at ECC. Rather, the data could be used in qualitative

assessments of contamination and in determining locations where further testing would be needed.

Historical sampling and testing information for ECC is discussed under the following headings:

- o Onsite surface water and sediment
- o Offsite surface water and sediment
- o Groundwater
- o Residential well water
- o Soil
- o Aquatic biota

#### ONSITE SURFACE WATER AND SEDIMENT

## Sampling and Testing

Table 3-1 summarizes the known surface water and sediment sampling events that took place onsite at ECC before RI activities began. Three general locations have been sampled: the cooling water pond, the north drum storage area pond, and the south drum storage area pond.

Sampling and testing procedures were not available for any of the events listed. However, all EPA samples were analyzed by labs selected and certified as part of the Contract Laboratory Program (CLP). Standard procedures are utilized

Table 3-1 HISTORICAL ONSITE SURFACE WATER AND SEDIMENT SAMPLING ECC SITE

Sampler	Sampling Date		Document Number	Sampling Location	No. of Water	Samples Sediment	Parameters Analyzed	Data Summary
ISBH	3/2/79	Water Laboratory, ISBH	24	Cooling water pond	1		COD, Pb, Hg, oil, phenol	Table 3-2
ISBH	6/8/79	Water Laboratory, ISBH	23	Cooling water pond; south storage area	2		As, Cd, Cr, Pb, Hg, Ni, Zn, oil, phenol, Cn	Table 3-2
ISBH	8/2/79	Water Laboratory, ISBH	33	Cooling water pond; south storage area	1		011, BOD, COD, Pb, N1, Zn	Table 3-2
ISBH	11/2/79	Water Laboratory, ISBN	35	Cooling water pond; north and south atorage areas	5		As, Cd, Cr, Pb, Hg, Ni, Zn, oil, phenol, pH	Table 3-2
ISBH	4/3/80	Water Laboratory, ISBH & Industrial Hygiene Laboratory	45	South storage area	1		PCB, Cd, Cr, Ni, Pb, Zn, Cu, phenol	Table 3-2
EPA	4/10/80	CLP <sup>a</sup> ; W. Coast Technical Service, Inc.	. 47	Cooling water pond; south storage area	2		Organic priority pollutants	Table 3-3
ISBH	4/17/80	Water Laboratory, ISBH	48	North and south storage areas	2		As, Cd, Cr, COD, Cu, Pb, N1, pH, phenol, 2n	Table 3-2
ISBH	3/10/81	Water Laboratory, ISBH	113	Cooling water pond	1	1	Metals, PCB's, volatile organics, others	Table 3-3
ISBH	4/29/81	Water Laboratory, ISBH	104	South storage area	2		Phenol, TOC, oil, volatile organics	Table 3-2
EPA	8/9/82	CLP	181	Cooling water pond	1		Organic priority pollutants	Table 3-3
EPA	10/18/82	CLP	209	Cooling water pond; north and south storage areas	4	1	Organic and inorganic priority pollutants	Table 3-3

a CLP = Contract Laboratory Program

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by these labs for the analysis of organic and inorganic priority pollutants.

All of the ISBH samples were analyzed by the ISBH Water Laboratory. The lab analyzed blanks and surrogate spikes with each set of samples. Duplicates were only occasionally analyzed.

## Results

Analytical results are summarized in Tables 3-2 and 3-3.

Table 3-2 presents the data for samples upon which only a limited analysis was performed. Table 3-3 summarizes the data for samples exposed to a more extensive analytical testing program.

The following inorganic chemicals were detected in the cooling water pond water samples at levels above EPA Water quality criteria:

- o Cadmium
- o Lead
- o Mercury
- o Nickel

A sample of the surficial oil layer from the north storage area pond taken on November 2, 1979, was found to contain

Table 3-2 HISTORICAL ONSITE SURFACE WATER SAMPLING RESULTS (ug/L) ECC SITE

														EPA Water
Water Quality	7	Cooling Wa	ter Pond			Sc	outh Drum Sto	rage Area Po	ondbox		North D	rum Storage A	rea Pond	Quality
Parameter	03/02/79	06/08/79	08/02/79	11/02/79	06/08/79	11/02/79	11/02/79	04/03/80	04/17/80	04/29/81	11/02/79	11/02/791	04/17/80	Criteria
_		_			_	_							_	0.022,c,d
yrsenic		4		11	1	6	4		18		60	900	7	
Cadmium		< 20		< 10	< 10	40	160	70	38		10	300	17	100
Chromium		390		< 10	1,100	40	250	770	380		1.6	104,000	1,000	10 <sup>b</sup> 50 <sup>b</sup> ,e
Lead	31,000	520	80	< 20	80	90	80	110	40		0.3	66,000	310	50, <sup>D</sup>
Mercury	< 10,000			< 0.5		< 0.5	< 0.5				0.9	< 200		0.144 <sup>D</sup>
Nickel		230	70	40	40	50	120	160	140		90	500	30	13.4 <sup>b</sup>
Zinc		580	290	150	2,300	140	260	290	90		1,090	18,000	3,100	NCA
		380	290	150	2,300	140	260				1,090	10,000	-	
Copper								460	838				11,100	NCA b
Phenol	8,800			65,300	28,000	22,500	25,500	22,400	13,000	10,000	35	3,000,000	8,900	3,500
011	80,000,000	18,000,000	8,300	20,000	110,000	180,000	63,000			62,400	3,032,000			
рĦ				6.3	2.0	7.3	7.2		6.9		7.1		7.1	
BOD			1,800,000											
	26,000,000		-,,						5,700,000				430,000,000	
TOC	20,000,000		6,000,000						2,.20,000	910,000			,,	
			0,000,000							310,000				0.00079 <sup>C</sup>
PCB								3.5						0.00079

NCA = Insufficient data available upon which to derive a criterion.

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Blank indicates parameter not analyzed.

a For the protection of human health assuming a daily ingestion of 2 liters of water.

For the protection of numeric absuming a contoxicity criteria.

Carcinogenicity criteria at the 10<sup>-5</sup> risk level.

Criteria applies to total trivalent arsenic.

Criteria applies to total hexavalent chromium.

f Oil layer.

Table 3-3 HISTORICAL ONSITE SURFACE WATER AND SEDIMENT SAMPLING RESULTS (ug/L) ECC SITE

				a.	SIIL					
			Cooling Water Po	nd		South Dr	um Storage Area	Ponds	North Drum	EPA Water
Organic					Sediment				Storage Area Pond	Quality
Priority Pollutants	04/10/80	03/10/81	08/09/82	10/18/82	03/10/81	04/10/80	<u>04/29/81<sup>f</sup></u>	10/18/82	10/18/82	Criteria a
1,1,-Dichloroethane	ND	4.4	17	ND	70	ND	< 5	ND	ND	NCA <sub>E</sub>
1,1,1-Trichloroethane	6,821	< 900	831	1,322	730	ND	160	621	1,266	18,400 <sup>b</sup>
1,1,2-Trichloroethane	16		< 2.8			ND	< 5			6.0
1,1-Dichloroethene	152	< 300	95	ND		NID	< 5	ND	ND	0.33 <sup>c</sup>
1,2-Dichloroethene	259	< 50	2,022	2,848	230	48		1,541	2,766	NCA
Tetrachloroethene	1,297	190	12	0.6	< 100	ND	260	1,176	71	8 <sup>C</sup>
Trichloroethene	3,873	< 600	191	673	470	ND	320	1,176	1,398	27°C
Methylene Chloride	5,470	240	1,329	3,908	1,500	485	180	3,873	5,548	1.9
Chloroform	ND	59	21	ND	90	< 10	9.1	ND	- <b>MD</b>	1.9
Trichlorofluoromethane	ND		< 2.7			14	< 5			1.9 <sup>C</sup>
Toluene	2,700	4,100			630	935	600,000			14,300 <sup>b</sup>
Nitrophenol	270		< 59			ND				NCA_
Pentachlorophenol	38		< 170			103		5	ND	1,010 <sup>b</sup>
Phenol	1,930	1,200	15,000	396	< 200	NID		460	325	3,500 <sup>b</sup>
2,4-Dimethylphenol	ND		260	251		349		236	121	NCA
2,4,6-Trichlorophenol	ND		< 62	5		ND		4	3	12 <sup>c</sup>
Benzene	ND	< 300	< 0.5	ND	90	ND	< 8	ND	463	6.6 <sup>c</sup>
Methylbenzene	ND		858	974		ND		1,035	1,132	,
Ethylbenzene	ND	600	110	ND	330	1,188	310	ND	ND	1,400 <sup>b</sup>
1,3-Dimethylbenzene	ND		98	ND		ND		ND	ND	
1,2 & 1,4-Dimethylbenzene	MD		79	ND		ND		ND	MD	<sub>b</sub>
1,3-Dichlorobenzene	ND		< 25	0.5		ND		17	92	400 <sup>b</sup>
1,4-Dichlorobenzene	ND		< 22	0.4		ND		15	86	400 <sup>b</sup>
1,2-Dichlorobenzene	ND		< 25	0.5		27		18	97	400 <sup>b</sup>
Diethylphthlate	27		86	47		433		32	ND	350,000 <sup>b</sup>
Dimethylphthlate	311		240	175		513		169	164	313,000
Butylbenzylphthalate	ND		< 290	1,122		ND		3,277	2,457	NCA
Di-n-butylphthalate	< 10		76	29		< 10		87	135	34,000
Napthalene	ND		< 23	12		ND		16	29	NCA
Isophorone	ND		3,200	ND		ND		ND	ND	5,200 <sup>b</sup>
P-Chloro-M-Cresol	ND				2,600	91				NCA
PCB's		< 50								0.00079 <sup>C</sup>

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Table 3-3 (Continued)

			Cooling Water P	ondbro	South Dr	um Storage Area	North Drum	EPA Water		
Organic Priority Pollutants	04/10/80	03/10/81	08/09/82	10/18/82	Sediment 03/10/81	04/10/80	04/29/81 <sup>f</sup>	10/18/82	Storage Area Pond 10/18/82	Quality Criteria
Arsenic		4.7		6.0	10,000			5.9	5.7	0.022 <sup>c,d</sup>
Cadmium		12		3.07				5.59	9.81	10
Chromium		150		286	19,000			326	320	50 <sup>b</sup> ,e 50 <sup>b</sup>
Lead		120		< 70	14,000			96.0	179	50 <sup>D</sup>
Mercury		0.2		< 0.1	30					0.144
Nickel		30		184	18,000			201	169	13.4 <sup>b</sup>
2inc		390		397	54,000			956	1,510	NCA
Copper		300		29.8	26,000			72.3	124	NCA
Aluminum		900		1,190	10,000,000			2,770	3,030	
Barium				138				172	183	
Beryllium		< 10		< 1	700			< 1	< 1	0.068 <sup>C</sup>
Cobalt				13.6				25.7	34.3	
Iron				6,840				14,600	19,800	
Manganese				2,370				2,370	1,960	
Boron				712				684	389	
Vanadium				8.6				13.3	12.6	
Silver				< 3				< 3	< 3 ′	50 <sup>b</sup>
Antimony				2.2				< 2	< 2	146
Thallium				< 2				< 2	< 2	13 <sup>b</sup>
Tin				< 40				< 40	62.6	
Ammonia		200		5,290	< 100					
Cyanide		52		16	< 625					200 <sup>b</sup>

ND = Not Dectected.

NCA = Insufficient data available upon which to derive a criterion.

Blank indicates parameter not analyzed.

b Toxicity criteria.

C Carcinogenicity criteria at the 10<sup>-5</sup> risk level.

Criteria applies to total trivalent arsenic.

Criteria applies to toal hexavalent chromium.

Oil layer.

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 $<sup>\</sup>stackrel{-}{a}$  Indicates no criteria is available. For the protection of human health assuming a daily ingestion of 2 liters of water.

arsenic, cadmium, chromium, lead, nickel, and zinc far in excess of the levels found for the pond water samples.

Listed in Table 3-3 are the organic priority pollutants found in at least one of the pond water samples above the detection limits. Background levels for these compounds are generally < 1 ug/L. The following eleven substances were found in the pond water samples at levels above EPA water quality criteria:

- o 1, 1, 2 Trichloroethane
- o 1, 1, Dichloroethene
- o Tetrachloroethene
- o Trichloroethene
- o Methylene chloride
- o Chloroform
- o Trichlorofluoromethane
- o Toluene
- o Phenol
- o Benzene
- o PCB's

Each of the onsite surface water areas sampled were found to contain levels of organic priority pollutants exceeding EPA water quality criteria.

One sample of the cocling water pond sediment was tested by the EPA. Inorganic pollutants reported in levels above background levels in sediment were arsenic, aluminum, chromium, nickel and copper. Organic pollutants reported in levels above background were 1, 1, dichloroethane, 1, 1, 1, trichloroethane, 1, 1, dichloroethene, trichloroethene, tetrachloroethene, methylene chloride, chloroform, toluene, benzene, ethylbenzene and PCB's.

#### OFFSITE SURFACE WATER AND SEDIMENT

# Sampling and Testing

Table 3-4 summarizes offsite surface water and sediment sampling episodes at ECC. The majority of sampling has been performed by the ISBH. The U.S. EPA performed one sampling episode. The United States Geologic Survey (USGS) performed three sampling episodes, collecting a total of 7 water samples and 15 sediment samples.

Sampling and testing procedure documentation was not available for the ISBH or EPA data. Testing procedures are known only in the general sense described earlier. Sampling and testing procedures employed by the USGS along with complete analytical results are described in: "Water and Streambed Material Data, Eagle Creek Watershed, Indiana, August 1980 and October and December 1982," Open File Report 83-215.

Table 3-4
HISTORICAL OFFSITE SURFACE WATER AND SEDIMENT SAMPLING
ECC SITE

	Sampling		Document		No. of	Samples		Data
Sampler	Date	Analytical Laboratory	Number	Sampling Location	Water	Sediment	Chemicals Analyzed	Summary
John Bankert	9/15/76	O.A. Laboratories	19	Creek	1		pH, COD, Fe, Cr, Ni, Pb, 2n, Cd, Cl	None
ISBH	6/8/79	Water Laboratory, ISBH	23	ε	3		As, Cd, Cr, Pb, Hg, Ni, oil, pH, phenol, Zn, PCB	Table 3-5
ISBH	7/31/79	Water Laboratory, ISBH	33	Finley Cr, Unnamed Ditch, Eagle Creek	5		011	None
ISBH	8/2/79	Water Laboratory, 1SBH	33	E, F	2		Oil, BOD, COD, Pb, Ni, 7n	Table 3-5
ISBII	11/2/79	Water Laboratory, ISBH	35	E, K	2		As, Cd, Cr, Pb, Hg, NI, oil, pH, phenol, Zn	Table 3-5
EPA	4/10/80	CLP - W. Coast Technical Services, In	nc. 47	E, J, K	3		Organic priority pollutants	Table 3-6
ISBH	4/17/80	Water Laboratory, ISBN	48	С, G, H, K	4		As, Cd, Cr, Cu, Pb, Ni, Zn, COD, pH, phenol	Table 3-5
1 SBH	8/25/80	Water Laboratory, ISBH	65A	A, B, L, M	4		PCB, As, Cu, Ph, Zn, diazinon	Table 3-5
USGS	8/25/80	USGS Laboratory	240	A, C, O, P		11	Metals, posticides, PCB, others	Table 3-7
ISBH	3/10/81	Water Laboratory, ISBH	113	A, C, E, N, P, Q, R	13	14	Metals, pesticides, PCB, volatile organics, others	Tables 3-6&7
ISBH	9/4/81	Water Laboratory, ISBN	137	B, E, H, I	4		011	None
ISBH	10/30/81	Water Laboratory, ISBH	149	D	1		Organic priority pollutants	Table 3-6
USGS	10/26/82	USGS Laboratory	240	A, P, S	4	4	Organic and inorganic priority pollutants	Tables 3-687
uscs	12/14/82	USGS Laboratory	240	A, S	3		Organic and inorganic priority pollutants	Table 3-6

a See Figures 3 and 4 for sample locations. Sampling location unknown.

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#### Results

Analytical results for the offsite surface water samples are summarized in Tables 3-5 and 3-6. Figure 3-4 indicates sampling locations. Table 3-5 presents data for surface water samples where only a limited analysis was performed.

Table 3-6 summarizes data for samples where more extensive analysis was performed. Data are presented for only those water quality parameters that had reported levels higher than upstream levels for at least one location.

Two inorganic chemicals were detected in offsite surface waters above EPA water quality criteria levels. Lead was found at sampling location B (downstream of the confluence of the unnamed ditch and Finley Creek) at 80 ug/L and at sample location Q (a small tributary to the unnamed ditch south of the landfill drive) at 250 ug/L. Nickel was reported at 20 ug/L at sample locations E (in the unnamed ditch alongside ECC) and K (upstream of ECC in the unnamed ditch).

These inorganic chemicals may be originating from ECC or NSL. Nearly all sample locations downstream of ECC and NSL showed at least one inorganic chemical at levels above the upstream values.

Table 3-5 HISTORICAL OFFSITE SURFACE WATER SAMPLING RESULTS (ug/L) ECC SITE

			S	AMPLE LOCATIO	NS DOWNSTREAM	OF ECC				SJ	MPLE LOCATIO	NS UPSTREAM O	F ECC	EPA Water
Water Quality	λ.	В	С		E		P	G	Н		(	L	М	Quality
Parameter	08/25/80	08/25/80	04/17/80	06/08/79	08/02/79	11/02/79	08/02/79	04/17/80	04/17/80	11/02/79	04/17/80	08/25/80	08/25/80	Criteria
Arsenic	1	3	3	4		3		18	1	1	1	2	ND	0.022 <sup>c,d</sup>
Cadmium			2	< 10		< 10		< 2	< 2		< 2			100
Chromium	10	60	160	< 10		< 10		< 10	< 10	< 10	< 10	13	10	50 <sup>b</sup> ,e
Lead	50	80	20	< 20	< 20	20	< 20	< 20	< 20	< 20	< 20	30	20	50 <sup>b</sup>
Mercury				< 0.1		< 0.1				< 0.1				0.144 <sup>b</sup>
Nickel			10	20	< 20	20	< 20	10	< 10	20	< 10			13.4 <sup>b</sup>
Zinc	76	79	80	20	< 20	< 20	< 20	10	< 10	20	< 10	70	148	NCA
Copper			65					6	4		< 4			NCA_
Phenol			9,800	2,000		< 5		1,500	< 5	7	< 5			3,500 <sup>b</sup>
011				3,400	< 1	2,800	< 1			42,000				
рĦ			7.2	1.7		7.2		6.8	7.7	7.3	7.7			
BOD					22,000		22,000							
COD			1,500,000		46,000		40,000	1,600,000	17,000		9,000			
PCB	120	10		< 0.1			•	•	•		•	10	1	0.00079 <sup>C</sup>

ND = Not detected.

NCA = Insufficient data available upon which to derive a criterion.

Blank indicates parameter not analyzed.

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Toxicity criteria at the 10<sup>-5</sup> risk level.

Criteria applies to total trivalent arsenic.

Criteria applies to total hexavalent chromium.

Table 3-6
HISTORICAL OFFSITE SURFACE WATER SAMPLING RESULTS (ppb)
ECC SITE

	SAMPLE LOCATIONS DOWNSTREAM OF ECC											
		S		A		С	D		E	R	Q	
Water Quality Parameter	10/26/82	12/14/82	03/10/81	10/26/82	12/14/82	03/10/81	10/30/81	04/10/80	03/10/81	03/10/81	03/10/81	
Aluminum	480	100	100	300	100	100			200	100	12,000	
Arsenic	4	2	0.7	6	3	1.1			0.8	0.6	4.0	
Barium	200	200		400	100				•			
Copper	12	4	5	9	8	4		_	4	5	17	
Iron	890	340		3,600	420			•				
Lead	6	3	< 10	5	5	10			10	20	250	
Manganese	120	70		280	80							
Magnesium			116			116			100	112	924	
2inc	10	20	< 10	10	30	< 10			< 10	10	60	
Strontium		•	170			170			150	120	650	
COD			21			4			4	5	17	
1,1 Dichloroethene	< 1	< 1	< 1	< 1	140	< 3	< 5	ND	< 1	< 6	< 1	
1,1 Dichloroethane	< 1	< 1	1.9	220	< 1	26	6	ND	1.2	< 1	< 1	
1,2 Trans-dichloroethene	< 1	< 1	< 20	1,000	9	< 20	< 5	45	< 1	< 20	< 20	
Methylene Chloride	< 1	< 1	1.1	< 1	< 1	18	350	< 10	3.5	< 10	< 1	
Trichloroethene	< 1	2	4.4	670	23	33	10	122	1	< 12	< 12	
Tetrachloroethene	< 1	1	1.2	37	< 1	2	1.8	< 10	< 1	1.2	2	
Toluene	< 1	2	< 3	7	2	5	< 6	< 10	< 3	< 3	< 3	
1,1,1 Trichloroethane	< 1	< 1	5.9	510	< 1	30	570	NED	< 1	9.1	5.6	
Chloroform	< 1	< 1		< 1	< 1		11.5	< 10			< 6	
1,1,2 Trichloro-1,2,2-												
trifluoromethane	< 1	< 1	< 2	< 1	< 1	< 40	< 5	ND	< 10	54	< 2	
Methyl ethyl ketone			< 52			270	1,900	ND	210	< 26	< 26	
2,4 Dimethylphenol	< 1	< 1		12	< 1		< 10	ND				
Phenol	< 1	< 1	< 0.2	2,200	< 1	< 0.2	< 10	14	< 0.2	< 0.2	< 0.2	
Butyl benzl phthalate	< 1	< 1		11	< 1		< 100	ND				
Bis (2-chloroethyl) ether	< 1	< 1		43	< 1		< 10	ND				
1-2 Dichlorobenzene	< 1	< 1		57	< 1		< 10	< 10				
Diethyl phthalate	< 1	< 1		6	< 1		< 20	ND				
Dimethyl phthalate	< 1	< 1		16	< 1		< 20	ND				
Di-n-butyl phthalate	< 1	< 1		27	< 1		< 30	< 10				
Bis (2-ethylhexyl)phthalate	< 1	< 1	< 0.35	13	< 1	< 0.35	< 100	ND	< 0.35	< 0.35	< 0.35	
Isophorone	< 1	< 1		360	< 1			ND				
n-Nitrosodimethylamine	< 1	< 1		9	< 1		-	ND				

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	SAMPLE LOCATIONS UPSTREAM OF ECC							
	J	K	N		P	Quality		
Water Quality Parameter	04/10/80	04/10/80	03/10/81	03/10/80	10/26/82	Criteria		
Aluminum			100	100	80	,		
Arsenic			0.2	0.7	3	0.022 <sup>d,e</sup>		
Barium					200			
Copper			< 4	< 4	9	NCA		
Iron					530			
Lead			10	< 10 <sub>.</sub>	6	, 50°		
Manganese					110			
Magnesium			200	220				
Zinc			< 10	< 10	10	NCA		
Strontium			90	160				
COD			6	8				
1,1-Dichloroethene	ND	ND	< 1	< 1	< 1	0.33		
1,1-Dichloroethane	ND	NID	< 1	< 1	< 1	NCA		
1,2-Trans-dichloroethene	ND	ND	< 1	< 1	< 1	NCA,		
Methylene Chloride	< 10	< 10	1.3	< 1	< 1	1.9 <sub>d</sub>		
Trichloroethene	ND	ND	< 1	< 1	< 1	27		
Tetrachloroethene	ND	ND	< 1	< 1	5	8 <sup>d</sup>		
To luen <b>e</b>	ND	ND	< 3	< 3	3	14,300		
1,1,1-Trichloroethane	ND	NID	< 1	< 1	< 1	18,400 c		
Chlorofor.	< 10	< 10			· < 1	1.9ª		
1,1,2-Trichloro-1,2,2-								
trifluoromethane	ND	NTD	< 2	< 2	< 1			
Methyl ethyl ketone	ND	ND	< 26	< 26				
2,4-Dimethylphenol	ND	ND			< 1	NCA		
Phenol	ND	ND	< 0.2	< 0.2	< 1	3,500 <sup>C</sup>		
Butyl benzyl phthalate	< 10	ND			< 1	NCA		
Bis(2-chloroethyl)ether	ND	ND			< 1	0.3		
1-2-Dichlorobenzene	ND	NED			< 1	400		
Diethyl phthalate	< 10	< 10			< 1	350,000		
Dimethyl phthalate	NED	NID			< 1	313,000		
Di-n-butyl phthalate	< 10	ND			< 1	34,000°		
Bis(2-ethylhexyl)phthalate	< 10	< 10	< 0.35	< 0.35	< 1	15,000 c		
Isophorone	ND	ND			< 1	5,200°		
n-Nitrosodimethylamine	ND	ND			< 1			

ND = Not Detected

NCA = Insufficient data available upon which to derive a criterion.

Blank indicates parameter not analyzed.

Blank indicates parameter not analyzed.

Parameters listed are only those that vary substantially from upstream value. See Appendix A for complete results.

For the protection of human health assuming a daily ingestion of 2 liters of water, 1982.

Toxicity criteria.

Carcinogenicity criteria at the 10<sup>-5</sup> risk level.

Criteria applies to total trivalent arsenic.

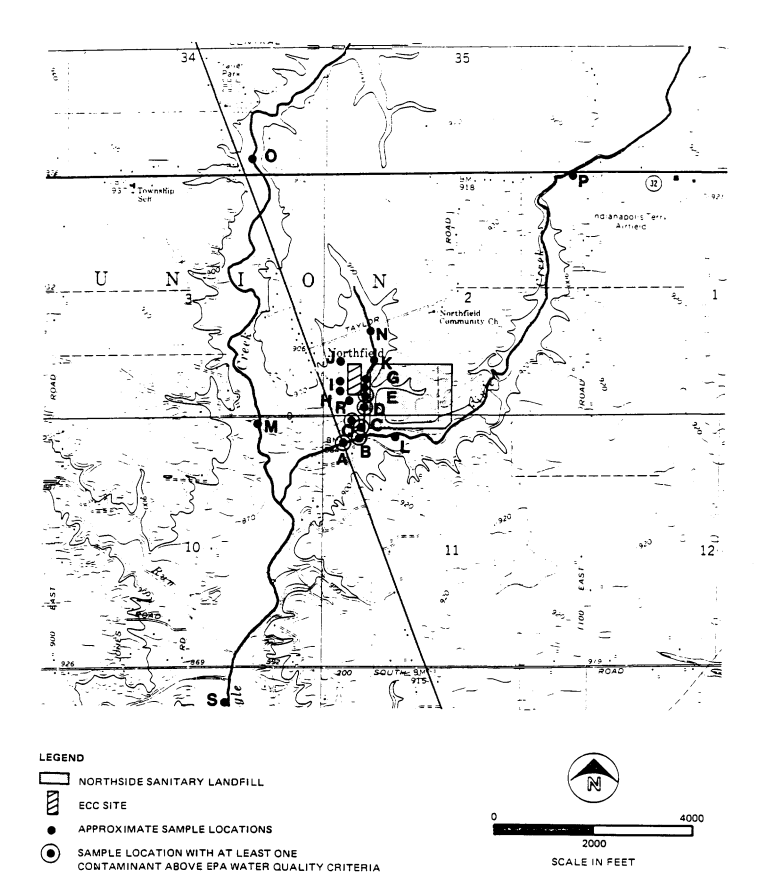


FIGURE 3-4
HISTORICAL SURFACE WATER
SAMPLE LOCATIONS
ECC RI REPORT

Eight organic priority pollutants were detected in surface water downstream of ECC at levels in excess of EPA water quality criteria. These pollutants, were:

- o 1,1 Dichloroethene
- o Methylene chloride
- o Trichloroethene
- o Tetrachloroethene
- o Chloroform
- o Bis (2-chloroethyl) ether
- o Phenol
- o PCB's

These were reported at sample locations A, B, C, D, and E (Figure 3-4).

Analytical results for surface water sediment samples are presented in Table 3-7. As with Table 3-6, this table only presents data for parameters that had at least one reported level greater than upstream values. Six compounds were reported at levels above upstream values: arsenic, chromium, copper, lead, DDD and PCB's.

#### GROUNDWATER

#### Sampling and Testing

Sampling and testing of groundwater from monitoring wells at GLT424/121 3-12

Table 3-7
HISTORICAL OFFSITE SURFACE WATER SEDIMENTS (ug/kg)
SAMPLING RESULTS
ECC SITE

Sediment				SAMPLI		SAMPLE LOCATIONS UPSTREAM OF ECC								
Quality	S		λ		(	3	E	Q	R	N	0		P	
Parameter a	10/26/82	08/25/80	03/10/81	10/26/82	08/25/80	03/10/81	03/10/81	03/10/81	03/10/81	03/10/81	08/25/80	08/25/80	03/10/81	10/26/82
Arsenic	< 1,000	1,000	5,700	1,000	3,000	4,400	10,000	5,200	8,800	6,500	< 1,000	2,000	6,600	1,000
Chromium	3,000	10,000	9,000	40,000	60,000	6,000	9,000	3,000	11,000	4,000	10,000	13,000	3,000	4,000
Copper	8,000	20,000	27,000	21,000	20,000	8,000	20,000	10,000	16,000	11,000	20,000	20,000	8,000	11,000
Lead	30,000	50,000	160,000	120,000	80,000	48,000	11,000	18,000	89,000	17,000	20,000	30,000	7,000	20,000
DDD	0.5	< 0.1		3.3	< 0.1						< 0.1	0.6		0.7
PCB's	5	120	< 1,000	72	10	< 1,000	< 1,000	< 0.5	< 1,000	< 1,000	1	10	< 1,000	13

a Sediment quality parameters listed are only those that vary substantially from upstream values.

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ECC is summarized in Table 3-8. Two monitoring wells were located onsite (Figure 3-5). Sampling has been performed by the ISBH on four occasions and by John Bankert on one occasion. Sampling results from the seven monitoring wells located along the perimeter of NSL are not summarized here.

Documentation of sampling and testing procedures was not found with any of the data. ISBH testing procedures are as described earlier. Testing procedures by O.A. Laboratories, Inc., laboratory for John Bankert, were not researched since only two samples were subjected to limited analyses.

#### Results

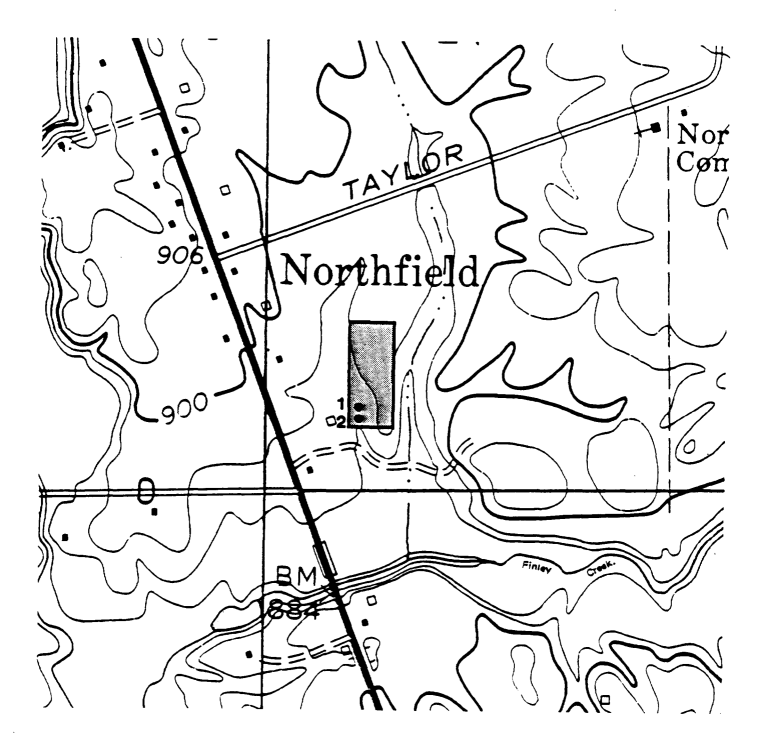
Analytical results are summarized in Table 3-9. Complete organic and inorganic priority pollutant analyses were not performed on any groundwater samples. For the samples tested, inorganic pollutants were not found at levels exceeding EPA water quality criteria. Two of the twelve organic priority pollutants were detected at levels above EPA water quality criteria. These were methylene chloride and trichloroethene. Other organic pollutants reported at levels above the detection limit were: 1, 2 - dichloroethane, 1, 1, dichloroethane, 1, 2 trans-dichloroethene, 1, 1, 1 trichloroethane, methyl ethyl ketone, toluene and isophorone.

Table 3-8
HISTORICAL GROUNDWATER SAMPLING
ECC SITE

Sample	Sampling r Date	Analytical Laboratory	Document Number	Monitoring Well Location	No. of Samples	Parameters Analyzed	Data Summary
John Ba	nkert 9/15/76	O.A. Laboratories	19	1, 2	2	pH, COD, Fe, Cr, Cr <sup>+6</sup> , Ni, Pb, Zn, Cd, C1	Table 3-9
ISBH	8/14/79	Water Laboratory, ISBH	29	1, 2	2	Cl, Fe, COD, TS, Hardness, Sulfates	None
ISBH	3/17/81	Water Laboratory, ISBH	86	1, 2	2	Metals, volatile organics, others	Table 3-9
ISBH	7/2/81	Water Laboratory, ISBH	121	1, 2	2	Metals, volatile organics, others	Table 3-9
ISBH	11/29/82	Water Laboratory, ISBH Hygrid Laboratory	243	2	2	Metals, volatile organics, others	Table 3-9

Well depths as follows: 1 = 71', 2 = 36'

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#### LEGEND

 APPROXIMATE MONITORING WELL LOCATION

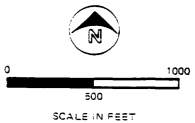


FIGURE 3-5
ECC MONITORING WELL
LOCATIONS (1982)
ECC RI REPORT

Table 3-9 HISTORICAL GROUNDWATER SAMPLING (ug/L) ECC SITE

		MONITOR WELL 1				HONITOR WELL 2	<del></del>	
Water Quality Parameter	09/15/76	03/17/81	07/02/81	09/15/76	03/17/81	07/02/81	01/29/82	01/29/82
luminum		< 100			100			
Arsenic		50	150		2.6	0.2	38	32
Barium			130			50		
Copper		< 4	< 4		18	< 4		
Throwium	< 100	< 10	15	< 100	< 10	< 10		
yanide		< 5			< 5			
Cadmium	< 100	< 2	< 2	< 100	< 2	< 2	< 2	< 2
Iron	2,600		2,000	32,000		< 50		
ead	< 100	< 10	< 10	< 100	< 10	< 10	< 10	10
lagnesium.		88,000			88,000			
ickel	< 100	< 10	< 10	< 100	< 10	< 10		
trontium		1,000			50			
inc	70	10	< 10	290	790	< 10		
OC .			3,900			2,100	28	31
OD	16,000	< 5,000	26,000	125,000	< 5,000	10,000	240	220
i (lab)	8.18	7.7	8.0	8.55		7.6	7.1	7.1
,2,-Dichloroethane		< 1	< 1		< 12	2.4	< 10	< 100
,1 Dichloroethane		< 1	< 1		50	41	160	130
,1 Dichloroethene			< 1		< 1	< 1	< 2	< 1
,2 Transdichloroethene		< 1	< 1		< 1	< 1	580	500
Methylene Chlorine		< 1	< 1		5.7	< 1	14	32
richloroethene		< 1	< 1		10	58	7.6	< 10
etrachloroethene		< 1	< 1		< 1	< 1	< 10	< 100
richlorofluoromethane		< 2	< 1				< 10	< 10
,1,1 Trichloroethane		< 1	< 1			1.2	30	< 100
hloroform			< 1			. < 1	< 10	< 100
1,1,2 Trichloro-1,2,2-tri-								
fluoromethane		< 2			< 2		ND	ND
ois(2-ethylhexyl)phthalate		< 350			< 350			
Methyl ethyl ketone		< 25	< 26		< 25	< 26	2,300	2,600

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		MONITOR WELL 1				MONITOR WELL 2			EPA Water Quality
Water Quality Parameter	09/15/76	03/17/81	07/02/81	09/15/76	03/17/81	07/02/81	01/29/82	01/29/82	Criteria
Phenol		< 200			< 200				3,500 <sup>b</sup>
Ethyl benzene			< 4			< 4	13	13	1,400
Toluene		< 4	< 4		< 4	5.5	13	15	14,300 <sup>L</sup>
Xylene		< 8	< 8	4	< 4	< 8	< 60	< 60	
Diazanon		< 0.3			< 0.3				
Isophorone							47	110	5,200
PCB		< 0.5			< 0.5				0.00079

NCA = Insufficient data available upon which to derive a criterion.

Blank indicates parameter not analyzed.

GLT424/33-2

a For the protection of human health assuming a daily ingestion of 2 liters of water, 1982.

Toxicity criteria.

Carcinogenicity criteria at the 10 risk level.

Criteria applies to total trivalent arsenic.

e Criteria applies to total hexavalent chromium.

# RESIDENTIAL WELL WATER

# Sampling and Testing

Residential well water sampling and testing activities are summarized in Table 3-10. Four sampling episodes were performed by the ISBH and one by Ira Jennings, a homeowner near ECC. Locations of the residential wells sampled are shown in Figure 3-6.

Documentation of sampling and testing procedures was not found with any of the data. ISBH testing procedures are as described earlier. Sampling of the Ira Jennings well was by Mr. Jennings. The sampling procedures used by him are unknown. Analysis of the sample was performed by Environmental Consultants, Inc. Testing and quality control procedures employed by the laboratory were not researched since only one sample was analyzed.

#### Results

Analytical results are summarized in Tables 3-11 and 3-12. Table 3-11 is a summary of residential well water sampling results for water quality parameters where levels above detection limits were reported. Table 3-12 is a list of additional organic pollutants analyzed by ISBH and not found above detection limits in any wells. Complete organic and

Table 3-10
HISTORICAL RESIDENTIAL WELL WATER SAMPLING
ECC SITE

Sampler	Sampling  Date	Analytical Laboratory	Document Number	Sampling Location	No. of Water Samples	Parmeters Analyzed	Data Summary
ISBH	8/14/79	Water Laboratory, ISBH	29	2	1	Cl, COD, Fe, Hardness, Sulfate	Table 3-11
ISBH	9/5/80	Water Laboratory, ISBH	71	3, 7, 9, 10, 13	5	Cd, Cr <sup>+6</sup> , COD, Cu, Fe, Pb, pH, phenol, TOC Hardness, Cl	Table 3-11
ISBH	3/5/81	Water Laboratory, ISBH	83	1, 2, 4, 5, 6, 7, 11, 12, 14	9	Metals, PCB, volatile organics, others	Tables 3-11,
Ira Jennings	6/26/82	Environmental Consultants, Inc.	241	8	1	Metals, methylene chloride, 1,1,2 trichloro ethane, tetrachloroethene	- Table 3-11
ISBH	12/9/82	Water Laboratory, ISBH	242	1	1	Volatile organics, others	None a

a No parameters with values above detection limits.

GLT424/32

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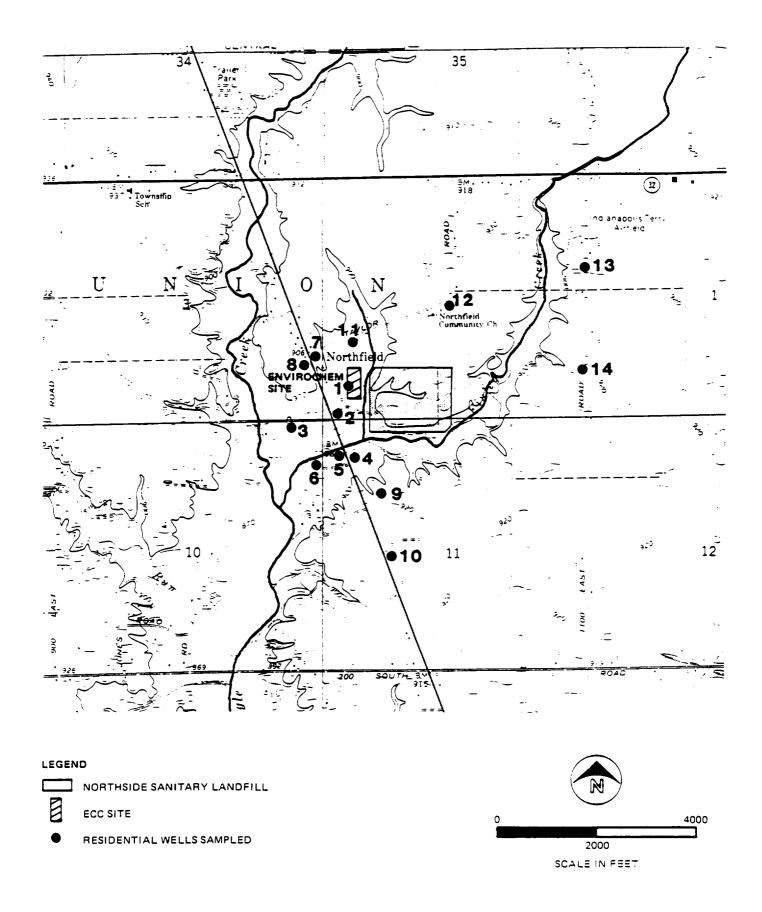


FIGURE 3-6
HISTORICAL RESIDENTIAL WELL
SAMPLING LOCATIONS
ECC RI REPORT

Table 3-11
HISTORICAL RESIDENTIAL WELL WATER SAMPLING RESULTS (ug/L)
ECC SITE

	1		!	3	4	5	6	7			9	10	11
Water Quality Parameter	03/05/81	08/04/79	03/05/81	09/05/80	03/05/81	03/05/81	03/05/81	09/05/80	03/05/81	06/26/82	09/05/80	09/05/80	03/05/81
Aluminum	< 100		< 100		< 100		< 100		< 100				< 100
Arsenic	0.9		0.8		< 0.2	< 0.2	0.3		3.1	10			0.4
Beryllium	< 10		< 10		< 10	< 10	< 10		< 10				< 10
Cadmium	< 2		< 2	< 2	< 2	< 2	< 2	< 2	< 2	9	< 2	< 2	< 2
Chromium-hex.	< 10		< 10	< 10	< 10	< 10	< 10	< 10	< 10		< 10	< 10	< 10
Chromium-tot.	< 10		< 10		< 10	< 10	< 10		< 10	< 3			< 10
Cyanide (free)	< 5		< 5		< 5	< 5	< 5		< 5				< 5
Iron	960	3,100	3,000	2,850	1,000	1,100	1,100	3,050	2,600		260	2,860	2,800
Lead	< 10		< 10	< 20	< 10	< 10	< 10	< 20	< 10	93	< 20		< 10
Mercury	< 0.1		< 0.1		< 0.1	< 0.1	< 0.1		< 0.1	< 0.5			< 0.1
Strontium	500		500	700	700	800		500				700	
Copper				11	< 4	< 4		< 4	< 4		26	< 4	6
Phenol				< 5				< 5			< 5	< 5	
Barium										403			
TOC				5,200				< 1,000			2,400	3,000	
COD		8,000		14,000				7,000			9,000	11,000	
Hardness (CaCo,)	272,000	332,000	356,000	248,000	268,000	272,000	272,000	424,000	432,000		224,000	288,000	348,000
Chlorides	< 5,000	7,000	10,000	< 5,000	< 5,000	< 5,000	< 5,000	16,000	15,000		6,000	5,000	7,000
pH (lab)	6.9		6.7	7.0	6.9	6.9	6.9	6.7	6.6		7.1	7.1	6.8
Methylene Chloride										20			
1,1,2 trichloroethane										31			
tetrachloroethene	< 1		< 1		< 1	< 1	< 1		< 1	46			< 1

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Water Quality Parameter	12 03/05/81	13 09/05/80	14 03/05/81	EPA Water Quality Criteria
Aluminum	< 100		< 100	
Arsenic	16		26	0.022 <sup>c,d</sup>
Beryllium	< 10		< 10	0.68
Cadmium	< 2	< 2	< 2	10.
Chromium-hex.	< 10	< 10	< 10	50 <sup>b</sup>
Chromium-tot.	< 10		< 10	170,000 b
Cyanide (free)	<b>&lt;</b> 5		< 5	200 <sup>b</sup>
Iron	3,900	1,030	2,300	
Lead	< 10	< 20	< 10	50, <sup>b</sup>
Mercury	< 0.1		< 0.1	0.144 <sup>D</sup>
Strontium	1,000		1,500	
Copper	< 4	< 4	< 4	NCA
Phenol		< 5		3,500 <sup>D</sup>
Barium				
TOC		5,500		
COD		14,000		
Hardness (CaCo,)	300,000	188,000	258,000	
Chlorides	9,000	< 5,000	< 5,000	
pH (lab)	6.9	7.3	6.9	
Methylene Chloride				1.9 <sup>c</sup>
1,1,2 trichloroethane				6.0
tetrachloroethene	< 1		< 1	8.0 <sup>C</sup>

NCA = Insufficient data available upon which to derive a criterion.

Blank indicates parameter not analyzed.

GLT424/34-2

For the protection of human health assuming a daily ingestion of 2 liters of water, 1982.

Toxicity criteria.

Carcinogenicity criteria at the 10 risk level.

Criteria applies to total trivalent arsenic.

# Table 3-12 RESIDENTIAL WELL WATER SAMPING ANALYSIS ORGANICS (ppb) ISBH SAMPLING 3/5/81

Parameter	Dectection Limit
Pyridine	< 1,000
Cresol	< 200
Heptaclor	< 0.02
Chloridane	< 0.24
Toluene	< 3
MIBK	< 12
Methyl ethyl ketone	< 26
Malathion	< 1.1
O-xylene	< 3
Benzene	< 3
1,1 dichloroethane	< 1
1,2 dichloroethene	< 1
trichlorofluoromethane	< 1
dichlorodifluoromethane	< 1
tetrachloroethene	< 1
trichloroethene	< 1
vinyl chloride	< 1
strobane	< 1
diazinon	< 0.3
dimethyl phenanthrene	< 500
trimethyl phenanthrene	< 500
PCB arochlor 1016	< 0.5
PCB arochlor 1242	< 0.5
PCB arochlor 1254	< 0.5
PCB arochlor 1260	< 0.5

 $<sup>^{\</sup>mathbf{a}}$  All nine residential well samples were reported to be below the detection limits for the parameters listed above.

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inorganic priority pollutant analyses were not performed on any well water samples prior to the onset of Superfund activities at the site.

The sample of the Ira Jennings well (well No. 8) was the only sample where a water quality parameter was detected at levels above the EPA water quality criteria. Lead, methylene chloride, 1, 1, 2 trichloroethane and tetrachloroethene were found to be above the EPA water quality criteria.

# SOIL

# Sampling and Testing

Sampling and testing of soil at ECC has been limited to one sample obtained by ISBH on March 2, 1979, from the dike between the cooling water pond and the unnamed ditch. Documentation of sampling and testing procedures was not found with the data.

#### Results

Analysis of the soil sample was limited to four parameters as follows:

o COD 30,000 ug/kg

o Pb < 1,000 ug/kg

o Hg 65,000 ug/kg

o Phenol 300 ug/kg

## AQUATIC BIOTA

# Sampling and Testing

Two studies, a bioaccumulation study on freshwater mussels and a biological assessment of stream ecosystems, have been performed in the vicinity of ECC. In the first study, the ISBH suspended live freshwater mussels, (Lampsilis radiata siluoides) in wire baskets at four locations on April 24, 1981, (Figure 3-7). On June 9, 1981 the mussels were taken out of the streams, wrapped in solvent-rinsed aluminum foil, and kept frozen until analyzed. Each sample consisted of five mussels.

The second study was performed by the Department of Zoology, Depauw University, from 1978 to 1980 as part of a larger biological monitoring program of fish populations and benthic macroinvertebrates. One of the watersheds studied was the Eagle Creek watershed, including Finley Creek. Figure 3-8 shows the locations of sample stations. Fish were collected using an electric seine. After being stunned, they were placed in live nets for later identification. Three passes

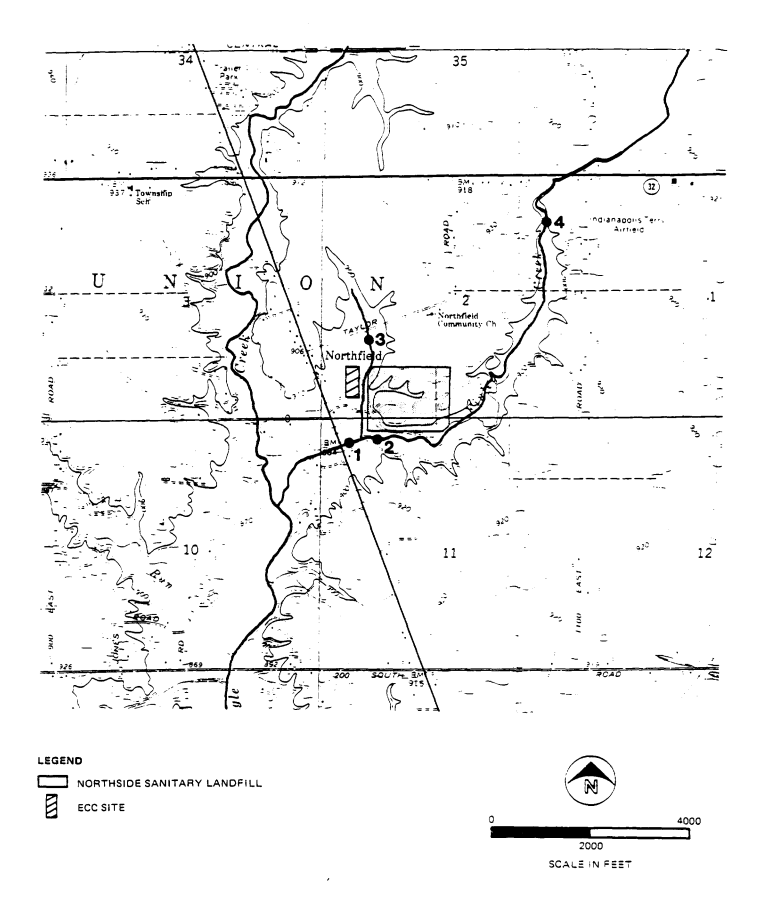
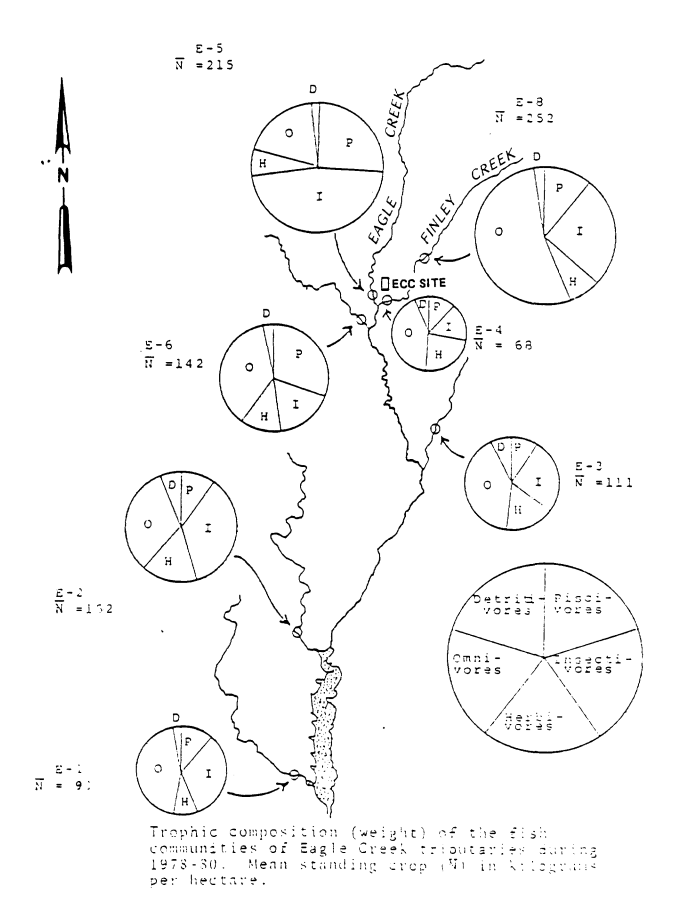


FIGURE 3-7
HISTORICAL BIOACCUMULATION
STUDY SITES
ECC RI REPORT



were made in each stream stretch. Benthic macroinvertebrates were collected with a square foot Surber sampler and a long handled dip net. Three replicates were collected at each station with each sampling device. Sampling normally took place once a month in May, June, July, August and October in 1978, 1979 and 1980. More complete sampling method descriptions are available in the report, "The Biological Monitoring Program of the Indiana MIP," by J.R. Gammon, M.D. Johnson, C.E. Mays and D.A. Schiappa.

#### Results

Analytical results from the mussel bioaccumulation study are presented in Table 3-13. The only parameter to be reported at levels higher downstream than upstream of ECC was arsenic.

Results of the Biological Monitoring Program assessment of fish population are shown in Figure 3-8. The mean standing crop of fish is much less at downstream station E4, compared to upstream station E8. Data on macroinvertebrates presented in the report is limited to a ranking of sample stations according to density, biomass or number of families (Table 3-14). Station E4 consistently ranked low in each category.

Table 3-13 FRESHWATER MUSSEL BIOACCUMULATION STUDY (ug/kg) ECC SITE

PARAMETER	SAMPLE LOCATION DOWNSTREAM OF ECC		SAMPLE LOCATIONS UPSTREAM OF ECC						
	1A	1B	2A	2B	3A	3B	<u>4A</u>	<u>4B</u>	
Fat (%)	51	51	58	60	41	57	87	98	
Arsenic	740	750	480	560	540	620	500	580	
Cadmium	300	340	260	320	320	300	220	280	
Chromium	400	400	< 200	600	400	200	300	1,000	
Copper	1,400	1,100	1,400	1,100	800	1,000	800	1,200	
Lead	< 800	< 800	< 800	< 800	< 800	< 800	< 800	< 800	
Mercury	< 30	< 30	< 300	< 200	< 300	< 200	< 300	< 200	1,000
Silver	< 100	< 100	< 100	< 100	< 100	< 100	< 100	< 100	
Aldrin	ND	ND	ND	ND	NID	ND	ND	ND	
Dieldrin	LOST	7	4	5	1	2	2	5	300
Chlordane	lost <sup>a</sup>	7	5	5	17	18	6	6	300
DDT	ND	ND	ND	ND	ND	ND	ND	ND	
Heptachlor	ND	ND	ND	ND	NID	ND	ND	ND	
Diazinon	ND	ND	ND	ND	NID	ND	ND	ND	
Strobane	ND	ND	ND	ND	NTD	ND	ND	ND	
Malathion	ND	ND	ND	ND	ND	ND	ND	ND	
PCB's	ND	ND	ND	ND	ND	ND	ND	ND	

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Sample Lost
b Federal Food and Drug Administration Action Level for substances in fish and shellfish

Table 3-14
MACROINVERTEBRATES
ECC SITE

RANK OF EAGLE CREEK STREAMS

Stream Mean Pool Depth	Fish (Composite Index)	Bivalvia (Density)	Tipulidae (Biomass)	Ephemeroptera (# of Families)	Baetidae (Density)
1. Mounts Run - E6	1. E5	1. E5	1. E5	1. E5	1. E5
2. Eagle (upper) - E5	2. E6	2. E2	2. E3	2. E6	2. E6
3. Fishback - E2	3. E2	3. E3	3. E2	3. E7	3. E7
4. Eagle (lower) - E7 <sup>b</sup>	4. E3	4. E7	4. E6	4. E2	4. E2
5. Little Eagle - E3	5. El	5. E6	5. E7	5. E3	5. E3
6. Finley - E4	6. E4	6. E4	6. El	6. E4	6. E4
7. School Branch - El	7.	7. El	7. E4	7. El	7. E1

a Invertebrates from Surber only.

#### Source:

The Biological Monitoring Program of the Indiana MIP. J.R. Gammon, M.D. Johnson, C.E. Mays and D.A. Schiappa. Department of Zoology, Depauw University.

GLT90/69

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No fish samples taken.

## PREVIOUS REMOVAL MEASURES

During March and April 1983, U.S. EPA removed and treated approximately 850,000 gallons of water from the cooling water pond to prevent overflows to the unnamed ditch.

Chemical Waste Management Inc. (Chem Waste) was hired by the U.S. EPA to conduct the ECC site surface cleanup. Chem Waste began onsite activities at ECC on July 11, 1983. On November 9, 1983, a Consent Decree was entered in U.S. District Court whereby some of the generators of waste sent to the site provided funding for completion of removal activities. Work under the Consent Decree was substantially completed on August 8, 1984. Tasks completed during this time period included:

- o Sampling and fingerprint testing of 29,192 drums.
- o Shipment offsite to a licensed hazardous waste disposal facility of 20,349 drums of waste.
- o Crushing onsite and shipment offsite to a licensed hazardous waste disposal facility of 9,558 empty drums.
- o Removal and shipment offsite to licensed disposal facilities of 282,500 gallons of liquids bulked from drums.

- o Removal and shipment offsite to licensed disposal facilities of 219,940 gallons of pumpable liquid hazardous wastes from the tanks (primarily flammable solvents).
- o Excavation and shipment offsite to licensed disposal facilities of about 5,200 yd3 of contaminated soil and cooling water pond sludge.
- o Removal and shipment offsite to a licensed hazar-dous waste treatment facility of about 4,500,000 gallons of contaminated cooling pond water.
- o Excavation and shipment offsite to a licensed disposal facility of 452 yd3 of contaminated soils from the polymer solidification pit.
- o Pressure washing of the concrete pad (about 27,000 ft<sup>2</sup>).
- o Cleaning of the processing building and equipment.

On August 1, 1984, U.S. EPA approved funding to undertake further surface cleanup work, some of which was reimbursed by the Consent Decree entered in November 1983. The following activities were completed:

- o Removal of remaining sludge from the bottom of the cooling water pond, and onsite containment.
- o Removal of remaining sludge from the bulk storage tanks.
- c Cleaning and/or disposal of the bulk tanks.
- o Removal of two underground tanks.
- o Removal of a leaking PCB-filled transformer.
- o Removal of miscellaneous piping.
- o Placement of a clay cover on the surface of the site, including filling in of the cooling water pond.

Remaining on the ECC site are some empty bulk tanks, the cleaned processing building with equipment, and additional areas of contaminated soils, including area beneath the concrete pad.

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#### Chapter 4

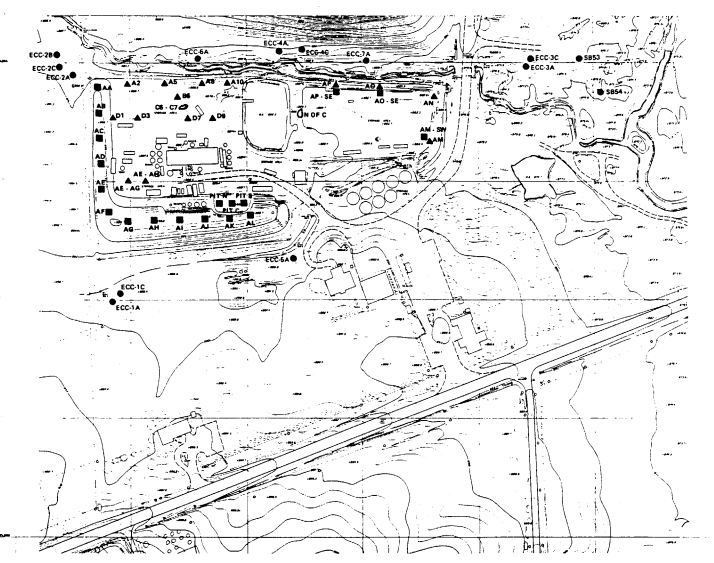
#### ANALYSIS OF SITE INVESTIGATIONS

# SOIL INVESTIGATIONS

# SCOPE AND METHODS

The purpose of the soil investigation was to collect data on the depth, areal extent and concentrations of hazardous constituents at potential contaminant source areas on the ECC site. An additional objective was to evaluate the dikes and embankments as possible sources of uncontaminated soil that could be used as cover material for potential remedial actions. A detailed summary of scope and methods is presented in TM 3-4 of Appendix A.

Soil sampling was performed in two phases, the first from May 7 through May 9, 1984, and the second, October 22 through October 26, 1984. In Phase 1, 18 surficial soil samples were taken along the north and west site embankments. Also, soil samples were collected from 2.5 foot deep soil borings with 2 inch diameter hand augers at 15 locations (Figure 4-1). Samples were screened for volatile organic contaminants (VOC's) using a field Organic Vapor Analyzer (OVA) and headspace analysis. The screening was used to select samples for the full CLP organic





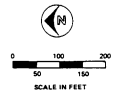


FIGURE 4-1
PHASE 1 SOIL BORING AND
SURFACE SAMPLE LOCATIONS
ECC RI REPORT

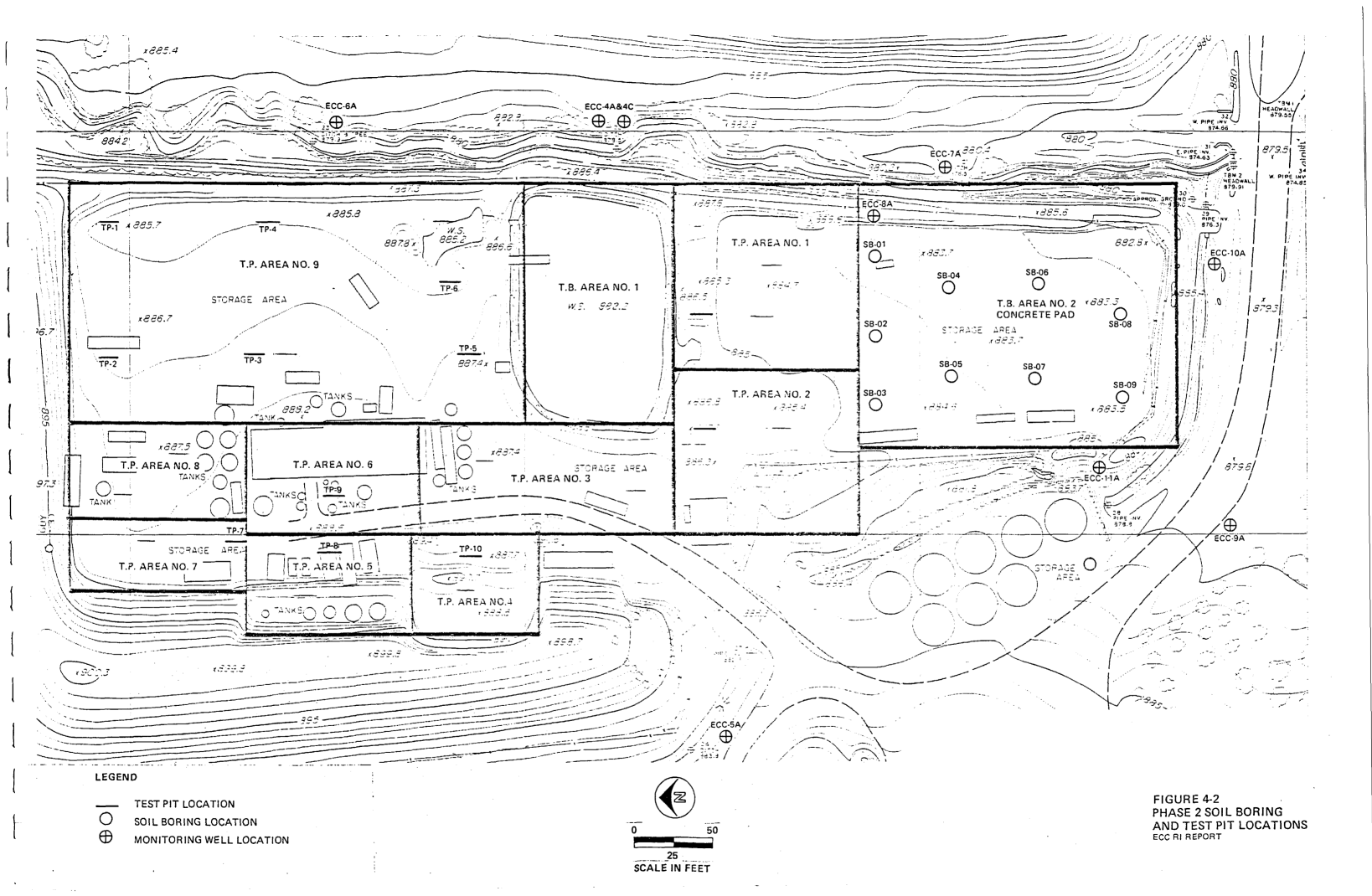
analysis. Site conditions were not favorable during Phase 1 sampling due to wet and muddy soils onsite to depths up to 2 feet. As a result the sampling results are considered indicative of contamination in the upper 2.5 feet of soil and no interpretation relative to variation of contaminants with depth is appropriate.

Phase 2 sampling consisted of nine soil borings to depths up to 12 feet through the concrete pad in the south area of the site and 12 test pits to depths up to 10 feet in the remaining areas of the site (Figure 4-2). Soil borings were advanced with a small drill rig and samples collected at 2 foot intervals with split spoons. Test pits were dug with a backhoe and samples collected at 2 foot intervals with hand augers. Samples were again screened in the field with an OVA and selected samples sent to the CLP for organic and inorganic analysis. Site conditions were more favorable than during Phase 1, although wet conditions did interfere with some of the sample efforts.

#### RESULTS

#### Inorganic Constituents

Only soil samples collected during the Phase 2 sampling were sent to the CLP for inorganic analysis. Tables 4-1 through 4-3 present the analytical results for these samples.



# TABLE 4-1 SOIL INDRGANIC RESULTS (mg/kg) TEST PITS SHOLOM DEPTH SAMPLES ECC RI Report

Sample Location: Depth (ft): Date Sampled: ITR Mumber:	TP-1 1-1.5 1 <del>0-22-84</del> NE4162	TP-2 1-1.5 1 <b>0-</b> 22-84 ME4164	TP-3 1-1.5 18-22-84 NE4165	TP-4 1-2 18-22-84 ME4166	TP-5 1-2 1 <del>0-</del> 22-84 NE4168	TP-5 2-3 1 <del>0-</del> 22-84 ME4169	TP-6 1-2 10-22-64 ME4170	TP-6 2-3 1 <del>8-</del> 22-84 ME4171	TP-7 1-2.5 1 <del>0</del> -23-84 HE4177	TP-8 1-2.5 10-24-84 NE4179	TP-9 1-3 10-24-84 ME4181	TP-10 1-3 10-24-84 ME4183	TP-11 1-2.5 18-24-84 ME4184	TP-12 0.5-3 10-24-84 ME4185
INERGRNIC COMPOUNDS														
ALININIM	6659	9990	44800	8000	4729	4878	8310	7189	4950	5639	3298	8310	10600	5988
ANTINONY ARSENIC BARIUM BERYLLIUM	7.1 [82] [ <b>9.</b> 6]	17 [73] [ <b>0.</b> 64]	5. 6 (28 <b>9</b> ) (3. 9)	(5, 9) (65) (8, 47)	9.7 (42)	16 [45] [ <b>0.</b> 37]	11 [82] [ <b>0,</b> 45]	7.4 157 <b>0</b> [1.4]	7.7 [81]	11 (51)	8. 6 [82] [8. 79]	[4, 8] [119] [8, 56]	6. 1 [69] [ <b>0.</b> 67]	8. 9 [49] [0. 44]
CADMILM CALCIUM CHROMILM COBALT COPPER	4.1 651 <b>80 +</b> 55 + [8.1] 38	7950 + 22 + [14] 38	126 <b>0000</b> 116 [51] 167	[2500] + 15 + [6, 5] [13]	191999 + 15 + [5, 1] 18	1 <b>03000 +</b> 12 + [6, 1] 17	23 <b>000 +</b> 93 <b>+</b> (12) 34	3.8 57 <b>800 +</b> 131 <b>+</b> [12] 77	93280 + 42 + [6.8] 31	11 <b>0000 +</b> 13 + (8.1) 21	4.5 50100 + 44 + (6.81 28	767 <b>88 +</b> 53 + [8.3] 39	3010 + 23 + (5.8) 25	194699 + 14 + [6.6] 28
IRON LEAD CYANIDE MAGNESILIN MINIGANESE	167 <b>08</b> 132 * 1.3 194 <b>00 *</b> 438	27 <b>986</b> 13 <b>*</b> 579 <b>0 *</b> 485	147 <b>000</b> 7.8 292 <b>000</b> 284 <b>0</b>	15300 11 + [2060] + 473	15000 9.1 28000 + 382	151 <b>00</b> 12 3 <b>0000 +</b> 327	15580 142 ± 0.86 8880 ± 299	188 <b>00</b> 393 * 111 <b>00</b> * 624 <b>0</b>	13400 135 + 2. 9 41500 + 366	16299 28 35199 + 371	11900 155 ± 19500 + 158	19308 189 ± 22460 ± 407	23600 11 3040 ± 109	17 <b>000</b> 8. 9 29900 + 324
MERCIRY NICKEL POTASSIUM SELENIUM SILVER	(20) (1290) (3, 8)	37 [1 <b>579</b> ]	[164] [1 <b>6589</b> ]	(121	[18] [116 <b>9</b> ]	[19] [1 <b>360</b> ]	[14] [1 <b>848</b> ]	[13] [9 <b>6</b> 5]	[5, 8) [2 <b>820</b> ]	[11] [11 <b>40</b> ]	(18) (1 <b>890</b> )	[22] [138 <b>8</b> ]	25 [1 <b>949</b> ]	[21] [141 <b>0</b> ]
SODIUM THALLIUM TIN VONADIUM ZINC	[21] [22] 121 *	[485] 32 90 +	[156 <b>89</b> ] [167] 477	[20] [22] 43 +	[1270] 25 [16] 48 +	[1630] [17] 56 +	[24] 164 +	[630] 33 517 #	(15) 232 +	[19] 73 <del>*</del>	[589] [24] [15] 122 #	[22] [24] 650 ±	35 82 +	[19] 59 +
PERCENT SOLIDS	78%	84%	90%	851	881	90%	88%	88%	841	87%	761	84%	985	98%

#### FOOTNOTES:

- E- Value is estimated or not reported due to the presence of interference.

  Duplicate analysis is not within control limits.

  Correlation coefficient for method of standard addition is less than 0.995.

  D- Positive values less than the contract required detection limit.

# TABLE 4-2 SOIL INDRSMIC RESULTS (mg/kg) TEST PITS INTERMEDIATE DEPTH SAMPLES ECC Site RI Report

Sample Location: Depth (ft): Date Sampled: ITR Number:	TP-1 4-5 18-22-84 NE4163	TP-4 2.5-3.5 18-22-84 HE4167	TP-6 4-5 10-22-84 ME4172	TP-7 2.5-4 10-23-84 ME4178	TP-8 2.5-4 10-24-84 HE4186	TP-9 3-5 10-24-84 NE4182	TP-10 3-5 10-24-84 HEA312	TP-11 3-5 18-24-84 NEA313	TP-12 3-5 19-24-84 MER314
INCREANIC COMPOUNDS									
ALIMINUM ANTIHONY	4628	13280	7928	5170	4670	5150	9970	5280	5848
ARSENIC BARIUM BERYLLIUM	(6, 1) (33)	29 137 [8.74]	[4.9] 173 <b>0</b> [1.5]	8. 4 [49]	[ <b>86</b> ] [2]	7.5 [47] [ <b>8.4</b> 3]	15 [63] [ <b>9.48</b> ]	[6. 0] [48]	6. 2 (46)
CADMILM CALCIUM CHROMIUM CUBALT COPPER	70100 + 13 + [7.1] 19	5868 + 25 + [13] 27	4.9 63 <b>899 +</b> 145 + [13] 85	92 <b>999</b> + 12 + (8.7) 19	27 87 <b>588 ±</b> 4 <b>0 ±</b> [9.4] 38	2.9 977 <b>60 :</b> 12 <b>:</b> [7.1] 18	3888 29 [11] 22	113 <b>999</b> 13 (8.5) 21	1 <b>9600</b> 15 [11] 29
IRON LEAD CYANIDE MAGNESIUM MANGANESE	14 <b>000</b> 8.5 238 <b>00 +</b> 352	31500 15 + 3740 + 780	20700 432 + 12300 + 6870	15600 54 6, 96 26780 + 479	145 <b>00</b> 142 + 4,4 253 <b>00</b> + 295	15800 15 27400 ± 379	221 <b>00</b> 12 3110 204 =	17486 7.7 27988 483 ±	16500 6.7 25700 389 #
MERCURY MICKEL POTASSIUM SELEMIUM SILVER	(17) (9 <b>35)</b>	36 (1 <b>840</b> ) (3, 8)	(15) (1 <b>030</b> )	[13] [1 <b>990</b> ]	[23] [1 <b>390</b> ]	[17] [1 <b>260</b> ]	[24] [1 <b>980</b> ]	[2 <del>8</del> ] [178 <b>9</b> ]	[19] [1500]
SODIUM THALLIUM	[1100]	••••••••	[480]		***********		[634]	[1560]	[1910]
TIN VONODIUM ZINC	(17) 53 +	36 90 *	37 570 +	[19] 62 +	[21] [17] 613 +	(17) 62 +	31 70	[19] 53	(28) 51
PERCENT SOLIDS	82%	81%	85x	89%	78%	93%	821	84%	89%

#### FOOTNOTES:

E- Value is estimated or not reported due to the presence of interference. 3- Duplicate analysis is not within control limits. 4- Correlation coefficient for method of standard addition is less than 0.995. []- Positive values less than the contract required detection limit.

TABLE 4-3 SDIL BORING INDROMNIC RESULTS (mg/kg) ECC Site RI Report

	INTERMEDIATE BORINGS											DEEP BORING	š		
Sample Location: Depth (ft): Date Sampled: ITR Number:	SB-01 2, 5-4 10-24-84 ME4186	SB-02 2.5-4 10-22-84 MEA310	SB-84 2-3, 5 10-24-64 MER320	SB-85 3-4.5 19-24-84 HEA325	SB-85 3-4.5 18-24-84 HEA324	SB-86 2-3.5 18-23-84 MEA318	SB-06 2.5-4 10-24-84 MEA317	SB-09 2.5-4 10-24-84 MEA316	SB-01 5.5-7 10-22-64 MEA309	SB-02 5.5-7 10-22-64 MEA311	SB-84 5-6.5 18-24-84 NEA319	SB-05 7.5-9 10-24-64 NEA323	SB-05 7.5-9 10-24-84 MEA322	SB-08 7-8.5 10-24-84 MEA321	58-09 5.7-7 10-24-84 MEA315
INDRGANIC COMPOUNDS															
ALINIMIN ARSENIC BARIUM BERYLLIUM CADMIUM	526 <b>0</b> (4.91 (35)	4588 8.6 [45]	6668 8.5 [54]	4650 18 [54] [_38]	514 <b>0</b> [4.6] [49]	5110 7.8 [35] [.36]	6548 7.3 [48] [.37] 4.4	5388 18 [32] [.38]	51 <b>00</b> 6.5 (81) 4.1	41 <b>88</b> 7.2 (35)	4378 [4.6] [38]	34 <b>88</b> [3, 7] [27]	339 <b>0</b> [4,5] [29]	4421 5.5 [4 <b>0</b> ]	6840 15 [44] [. 39]
CALCIUM CHROMIUM CUBALT COPPER IRON	11 <b>0000</b> 15 (51 23 16 <b>000</b>	192900 12 [11] 18 15399	108000 15 [10] 25 19800	121 <b>000</b> 13 [18] 21 1 <b>9200</b>	199 <b>000</b> 12 [9.6] 21 161 <b>90</b>	199000 13 [6, 6] 20 14400	194999 18 [11] 26 2 <b>9599</b>	113000 14 [9.5] 20 16400	1 <b>84889</b> 15 (8, 5) 18 151 <b>98</b>	187999 11 [6-6] 18 14389	1 <b>08000</b> 13 [9, 9] 23 1 <b>6400</b>	197969 9.6 [7.1] 19 13298	140000 10 [6, 8] 21 13800	119 <b>000</b> 9.8 [6.5] 18 151 <b>00</b>	68888 17 [6.5] 24 28788
LEAD NAGNESTUM MANGAMESE NICKEL POTASSTUM	7. 2 26499 289 [13] [1489]	9, 3 28600 • 344 [15] [1638]	9. 1 27300 451 23 [1750]	26 27 <b>098</b> • 4 <b>09</b> [19] [1 <b>550</b> ]	5.6 39499 4 314 [16] [1759]	8.3 33300 • 306 [18] [1640]	9 28799 481 24 (2938)	7.7 34100 # 316 # [13] [1450]	6.5 274 <b>88</b> 555 (291 (1490)	7.2 28988 334 15 [1628]	7.1 29500 337 [19] [1630]	4.5 24800 285 (133 (1240)	5 28789 485 [15] [1289]	7.1 39299 399 [16] [1599]	17 21386 390 (181 [1190]
SILVER SODIUM TIN VANADIUM ZINC	(859) (29) 51	[944] [16] 47	[1649] 39 [23] 69	[1 <b>090]</b> 19 (18] 54	[98 <b>0</b> ] 17 [2 <b>8</b> ] 66	(1290) (19) 55	[3, 3] [1 <b>486</b> ] [25] 68	(1390) (20) 56	[673] [19] 47	(958) (15) 56	[143 <b>0</b> ] [17] 44	[983] [16] 54	[11 <b>06</b> ] [15] [38]	[121 <b>8</b> ] (15] 41	[119 <b>6</b> ] [22] 65
PERCENT SOLIDS	98%	81%	887	92%	90%	90%	89%	91%	92%	98%	91%	93%	923	91%	84%

#### FOOTNOTES:

- E- Value is estimated or not reported due to the presence of interference.

  4- Duplicate analysis is not within control limits.

  4- Correlation coefficient for method of standard addition is less than 0.995.

  C1- Positive values less than the contract required detection limit.

Sampling locations are presented in Figures 4-1 (Phase 1 sampling) and 4-2 (Phase 2 sampling).

Background Concentrations. General standards are not established for inorganic metal concentrations in soil.

Therefore, metal concentrations reported for soil samples from the site are compared with typical concentration ranges and estimated background levels for these inorganic constituents to determine if contamination is present.

Onsite background inorganic concentrations were estimated using eight Phase 2 soil samples. Results of organic analysis indicated that organic contaminants were either not present or present only in relatively minor concentrations in these samples. Therefore, these eight samples were considered least affected by waste handling operations at the site and selected to estimate background levels.

For each inorganic constituent, the mean concentration, standard deviation, and the 95 percent confidence interval of the mean was calculated using the analytical results from the eight selected soil samples. These background values are presented in Table 4-4.

Also, shown in Table 4-4 are typical concentration ranges for inorganic constituents in soil. These published ranges

Table 4-4 TYPICAL AND BACKGROUND CONCENTRATIONS OF METALS IN SOIL (mg/kg)

			Onsite Background				
		Observed Range in Background Samples	Mean	Standard Deviation	Upper 95 percent Confidence Interval of the mean	Typical Range in Soil	Concentration Range in Soil
Aluminum	λl	4,100 - 10,600	6,151	2,594	12,290		10,000 - 300,000
Antimony	Sb	<25 - 42	-	-	-	2 - 10	0.2 - 150
Arsenic	λs	4.6 - 17	7.6	3.9	16.8	1 - 50	0.1 - 194
Barium	Ba	33 - 81	5.3	18.7	97.2	100 - 3,000	100 - 3,000
Beryllium	Be	<0.3 - 0.67	-	-	-	0.1 - 40	0.1 - 40
Cadmium	Cđ	<2 - 4.1	-	-	-	0.01 - 0.7	0.01 - 7
Chromium	Cr	11 - 15	13	2	17.7	1 - 1,000	5 - 3,000
Cobalt	Co	5.8 - 14	8.4	2.6	14.6	1 - 40	0.05 - 65
Copper	Cu	18 - 30	21.5	4.0	31.0	2 - 100	2 - 250
Cyanide	Cn	Less than 0.5	Less than 0.5	-	-		
Iron	Fe	14,000 - 27,000	17,950	4,754	29,190		100 - 550,000
Lead	Pb	6.7 - 15	9.5	3.1	16.8	2 - 200	<1 - 888
Manganese	Mn	109 - 555	369	131	679	20 - 3,000	20 - 18,300
Mercury	Hg	Less than 0.05	Less than 0.05	-	-	0.01 - 0.3	0.01 - 4.6
Nickel	N1	15 - 37	21.2	7.0	37.8	5 ~ 500	0.1 - 1,530
Selenium	Se	Less than 2.5	Less than 2.5	-	-	0.1 - 2.0	0.1 - 38
Silver	λg	Less than 2.5	Less than 2.5	-	_	0.01 ~ 5	0.01 - 8
Thallium	T1	Less than 3.0	Less than 3.0	-	-		0.1 - 0.8
Tin	Sn	Less than 14	Less than 14	-	-	2 - 200	1 - 200
Vanadium	V	17 - 35	21.4	7.6	39.4	20 - 500	3 - 500
Zinc	2n	44 - 90	60.9	16.6	100	10 - 300	1 - 2,000

#### Notes:

GLT360/69

a Onsite soil samples used to estimate background soil values are: SB01 (5.5-7 ft), SB02 (5.5-7 ft), SB04 (5-6.5 ft), TP-1 (1-1.5 ft), TP-1 (4-5 ft), TP-2 (1-2.5 ft), TP-9 (3-5 ft), TP-11 (3-5 ft).

Source: W.L. Linday, Chemical Equilibrium in Soils, 1979.

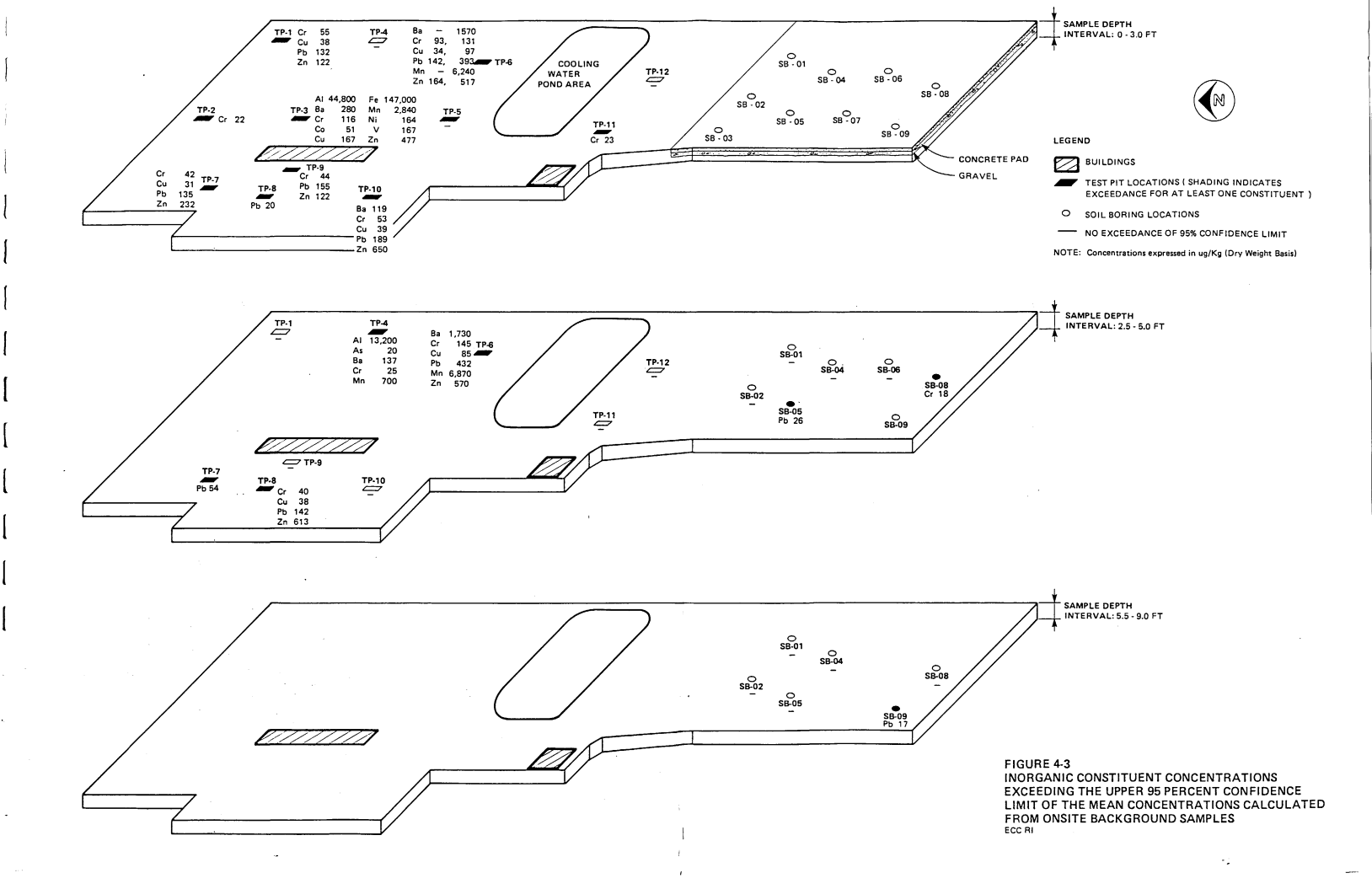
Sources: H.J. M. Bowen, Environmental Chemistry of the Elements, 1979; URE, A.M., et al., Environmental Chemistry, 1983; Parr, J.F., Marsh, P.B., KLa, J.M., Land Treatment of Hazardous Wastes, 1983.

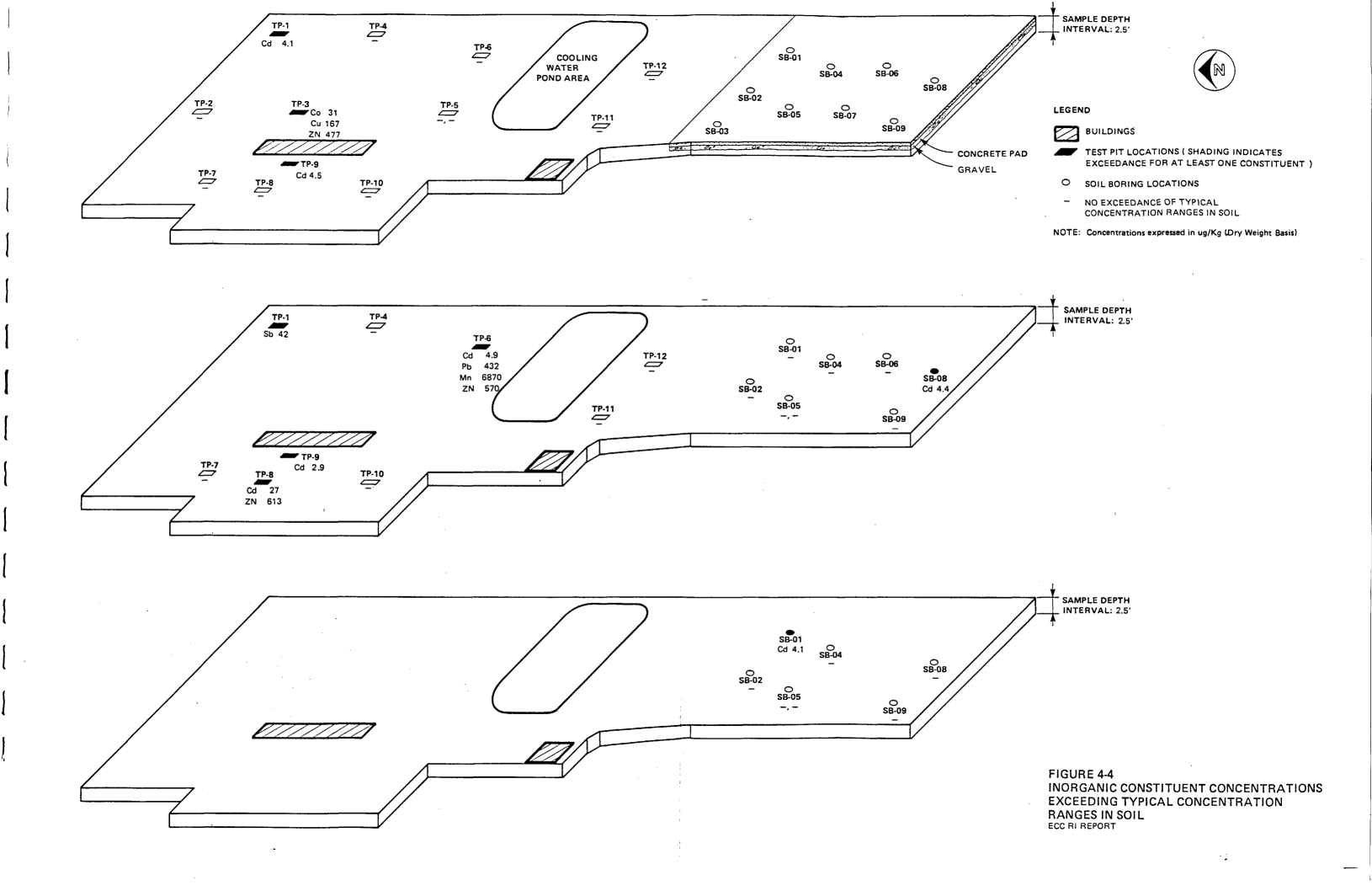
were developed from concentration measurements in soil sampled throughout the United States.

Inorganic Contamination. Inorganics most frequently exceeding the comparison criteria include cadmium, chromium, copper, lead, and zinc. Other less frequently exceeding inorganic constituents include aluminum, arsenic, barium, cobalt, iron, manganese, nickel, and vanadium. Figure 4-3 summarizes the distribution of inorganic constituents exceeding the upper 95 percent confidence limits of background concentrations. Figure 4-4 summarizes the distribution of inorganic constituents exceeding typical concentrations in soil.

Observations regarding the comparison of the inorganic analysis results with estimated onsite background values for soil are:

- o The largest variety of inorganics constituents exceeding background values are reported in shallow (0-3 feet) soil samples.
- o The number and frequency of inorganic constituents exceeding background values decreases with depth.





- o Inorganic constituents that represent the most widespread exceedance of background values are chromium, copper, lead, and zinc.
- o Inorganic constituent exceedance of background values in soil beneath the concrete pad is minor relative to the soil in the northern drum and tank storage areas.

Observations regarding the comparison of inorganic analysis results with typical ranges for soil are:

- o Only antimony, cadmium, cobalt, copper, lead, manganese, and zinc were reported in soil samples at concentrations exceeding the typical range in soil.
- o Only cadmium, lead, and zinc were reported in more than one sample at concentrations exceeding the typical range in soil.
- o Inorganic constituent exceedance of the typical ranges in soil for samples beneath the concrete pad is minor relative to the soil in the northern drum and tank storage areas.

## Organic Compounds

Soil samples collected during the Phase 1 and 2 sampling activities were analyzed for volatile organics, acid extractable, base/neutral extractable, pesticide, and PCB compounds using the CLP. Analytical results are presented in Tables 4-5 through 4-8.

Background Concentrations. General standards are not established for organic compound concentrations in soil. Therefore, organic compound concentrations reported for soil samples from the site are compared with background concentrations to determine if contamination is present. Many of the organic compounds analyzed for during this RI are not naturally occurring compounds and their presence indicates the influence of man's activities on the soil. Also, analysis of several soil samples from the site did not detect any priority pollutant organic compounds or other organic compounds on the CLP's hazardous substances list. Therefore, this RI report considers the detection of organic compounds in soil samples analyzed for by the CLP's routine analytical services as evidence of contamination.

Organic Contamination, Phase 1 Sampling. Analysis of soil samples collected during Phase 1 sampling activities detected a wide variety of organic contaminants. Organic contaminants included volatile organic, acid extractable, base/neutral extractable, and pesticide compounds. PCB's were detected in only one Phase 1 soil sample.

#### TABLE 4-5 SOIL ORGANIC RESULTS (ug/kg) PHRSE I SAMPLING ECC Site RI Report

		N		OIL SAMPLE ORTHWEST E		<b>;</b>			SURFACE SI	DIL SAMPLE	S				SOIL BORIN	NG SAMPLES	
Sample Location: Depth (ft): Date Sampled: OTR Number:	AA 8-8.5 5-8-84 E-7244	AC 8-0.5 5-8-84 E-7245	AE 8-0.5 5-8-84 E-7246	AG 8-8.5 5-8-84 E-7247	AI <del>0-0</del> .5 5-8-84 E-7248	AK 9-0.5 5-8-84 E-7249	AL 9-8.5 5-8-84 E-7258	AM-SH 9-0.5 5-9-84 E-7255	AO-SE 0-0.5 5-8-84 E-7251	AP-SE 9-8.5 5-8-84 E-7252	N OF P 5-9-84 E-7253	N OF PO 5-9-84 E-7254	AN 8-0.5 5-9-84 E-7256	AE-AH 0-0.5 5-9-84 E-7257	AE-A6 0-0.5 5-9-84 E-7258	B-6 0-0.5 5-8-84 E-7259	D-7 1.5-2 5-8-84 E-7260
VOLATILE COMPOUNDS																	
1, 2-DICHLOROETHANE 1, 1, 1-TRICHLOROETHANE 1, 1-DICHLOROETHANE CHLOROFORM TRANS-1, 2-DICHLOROETHENE								676 <b>000</b>	289 17 <b>506</b> 580 797 <b>00</b>	19 <b>3500</b> 700 890 1500	74114 <b>00</b>	451 <b>9909</b>	40 60 20 100	48000	27 <b>8900</b>	1203200 41800 41800	635 <b>000</b> 176 <b>00</b>
CIS-1, 3-DICHLOROPROPENE ETHYLBENZENE ETHYLBUE CHLORIDE CALOROMETHANE	88 79	10	18	20	29	50	50	262 <b>989</b> 51 <b>5000</b>	6 <b>99</b> 24 <b>99</b>	2500	1212 <b>90</b> 1418 <b>90</b>	514 <b>999</b> 12 <b>9999</b>	10	9000 34000	12000 5649000 35000	155 <b>000</b> 655 <b>00</b>	12 <b>989</b> 0 94 <b>00</b> 0
TETRACHLOROETHENE								4116888	570	46 <b>00</b>	6172 <b>88</b>	625000		131 <b>999</b>	238000	639900	744100
TOLLENE TRICHLOROETHENE VINYL CHLORIDE ACETONE 2-BUTANONE							5	751000 K 4214000	14889 1889 6489 38389 5289	2888	6871 <b>00</b> 5 <b>888290</b>	674 <b>000</b> 2 <b>006000</b>	60	80000 147000	273 <b>999</b> 56 <b>4999</b>	470788 2135788 99288	964 <b>900</b> 1375 <b>900</b> 896 <b>00</b>
4-METHYL-2-PENTANONE STYRENE TOTAL XYLENES	*********	******	••••		••••••	• • • • • • • • • •		1160000	73 <b>0</b> 15 <b>000</b>	•••••	26 <b>88</b> 7 <b>07880</b>	K 2299 I	·······	5 <b>000</b> 97 <b>000</b>	19000 633000	76 <b>00</b> 8826 <b>00</b>	29600 13800 607000
TOTAL VOLATILES	150	19	10	20	20	50	52	11728400	175868	296490	15769380	8796200	290	551888	7793888	5733100	4689700
ACID COMPOUNDS  2, 4-DIMETHYLPHENOL PHENOL BENZOIC ACID 2-HETHYLPHENOL								36866 18866 93186	K 7299 11999	К	3676 <b>20</b> 613 <b>20</b>	88006 447000 K 142600	K 16 <b>00</b>	24599 K 28299 H 29899	138 <b>999</b> ( 28 <b>999</b>	11 <b>4000</b> K 13 <b>000</b>	119 <b>000</b>
4-HETHYLPHENOL								52900	10000		87988	К 535600		67800	36788	K 510000	31000
BASE/NEUTRAL COMPOUNDS		• • • • • • • • • • • • • • • • • • •	22.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2					199100	182 <b>90</b>		516800	1213299	16 <b>00</b>	141300	203600	754 <b>000</b>	173 <b>890</b> =======
1, 2, 4-TRICHLOROBENZENE 1, 2-DICHLOROBENZENE 1, 4-DICHLOROBENZENE 1, 2-DIPHENYLHYDRAZINE HEXACHLOROBUTADIENE								159 <b>89</b> 686 <b>89</b>	33700		389600 534100 576000	49 <b>000</b> 3337 <b>00</b>		84189	252786	216 <b>9999</b> 4 <b>999</b>	119000 172000 K
ISOPHORONE NAPHTHALENE NITROSENZENE NITROSENZENE N-NITROSODIMETHYLAMINE N-MITROSODIMETHYLAMINE		• • • • • • • • • • • • • • • • • • • •	**********	40	1400	40	• • • • • • • • • • • • • • • • • • • •	41900 38300		•••••	4 <b>09200</b> 2983 <b>00</b>	44808 55799		417 <b>00</b> 261 <b>00</b>	593 <b>00</b> 4 <b>08000</b>	340000 470000	122 <b>000</b> 99 <b>000</b>
N-NITROSODIPROPLYAMINE BIS(2-ETHYLHEXYL) PHTHALATE BENZYL BUTYL PHTHALATE DI-N-BUTYL PHTHALATE DI-N-OCTYL PHTHALATE	230	 48 К		49		•••••	• • • • • • • • • •	7552 <b>98</b> 1282 <b>989</b> 679 <b>98</b> 1278 <b>88</b>	12 <b>906</b> 48300 42500 8300	17890	774699 209999 K 78699	685900 366000 79000 K 84000	970	291908 K 85000 14300 8900	458109 268000 112200 22600	3800000 1000000 K 300000	226090 61000 11000 34000
DIETHYL PHTHALATE DINETHYL PHTHALATE PHENANTHRENE 2-NETHYLNAPHTHALENE	• ••••••	•••••	•••••			•••••		7200	К	••••••	104000	35000 44900		46 <b>08</b> 88 <b>00</b>	25 <b>489</b> K 551 <b>99</b>	130000	8 <b>999</b> 31 <b>999</b>
	248		) (	80	1480	40	*******	2396800	201470	17888	3359300	1777200	970	565400	1661400	8294889	883860

## TABLE 4-5 SOIL GREANIC RESULTS (ug/kg) PHASE I SAMPLING ECC Site RI Report

		N	Surface s Orth and N	OIL SAMPLE: ORTHWEST E	s from Mbankments	<b>.</b>			SURFACE S	OIL SAMPLE	S				SOIL BORI	NG SAMPLES	
OTR Number:	AA 9-8.5 5-8-84 E-7244	AC 8-8.5 5-8-84 E-7245	AE 8-8.5 5-8-84 E-7246	A6 9-0.5 5-8-84 E-7247	AI 9-0.5 5-8-84 E-7248	AK 9-0.5 5-8-84 E-7249	AL 9-0.5 5-8-84 E-7250	AM-SH 8-0.5 5-9-84 E-7255	A0-SE 9-0.5 5-8-84 E-7251	AP-SE 9-9.5 5-8-84 E-7252	N OF P 5-9-84 E-7253	N OF PD 5-9-84 E-7254	AN 0-0.5 5-9-84 E-7256	AE-AH 9-0.5 5-9-84 E-7257	AE-A6 8-8.5 5-9-84 E-7258	8-6 8-0.5 5-8-84 E-7259	D-7 1.5-2 5-8-84 E-7268
PESTICIDES																	
DELTA-BHC GAMMA-BHC (LINDONE) HEPTACHLOR ALDRIN ENDOSULFAN I										40	83 <del>00</del>	760	10	948	260 90 20	170 170 210	546
DIELDRIN 4, 4-DDE ENDRIN ENDOSULFAN II 4, 4-DDD	••••••	•••••	•••••	**********	•••••	18	••••••	450	• • • • • • • • • • •	140 190	29 838 1 <b>999</b> 9	28 729 63 <b>86</b>		1 <b>96</b> 679	110 110 1080	1649	70 1120 1110 590
ENDRIN ALDEHYDE ENDOSJUFAN SJUFATE 4, 4-DDT WETHOLYCHLOR CHLORDANE TOXAPHENE	78			*******	•••••	• • • • • • • • • •	••••••	*********	•••••	500	12199 4999 28999 2799 19899	9400 3300 21000 2300	49	1389	2800	3200	2000 1900 3600
TOTAL PESTICIDES	78	0	9	0	0	18	9	450	0	870	77650	438 <b>00</b>	50	2160	4470	3910	18444
PCB¹ s																	
AROCHLOR-1016 AROCHLOR-1232 AROCHLOR-1248											18888 16288 18888						
TOTAL PCB's	8	0	0	9	8	0	8	9	0	8	37800	9	*	0	9	9	
DIOXIN									<del>-</del>							•	_
2, 3, 7, 8-TETRACHLORODIBENZO-P-DIOXIN			*****		********					F# 2112222		25-1-2-2-22	B##########	.=====		7.6	6 =======
PERCENT MOISTURE	14.6×	14.1%	13.8%	11.3%	11.4%	12%	11.8%	16.9%	15.5%	14.2%	48.1%	38.5%	16.9%	13.5≭	15≱	29.27	21.

#### FOOTNOTES:

- A. Tentatively identified compound concentrations are estimated. A 1:1 response is assumed.

  B. Analyte has been found in the laboratory blank as well as the sample. Indicates probable contamination.

  C. Applies to pesticide parameters where the identification has been confirmed by BC/MS.

  J. Indicates an estimated value. When mass spectral data indicates the presence of a compound that meets the identification criteria and the result is less than the specified detection limit but greater than zero.

  K. Actual value, within the limitations of the method is less than the value given

## TABLE 4-6 SOIL ORGANIC RESULTS (ug/kg) TEST PITS SHALLON DEPTH SAMPLES ECC Site RI Report

Sample Location: Depth (ft): Date Sampled: OTR Number:	TP-1 1-1.5 18-22-84 E4901	TP-2 1-1.5 10-22-84 E4903	TP-3 1-1.5 19-22-84 E4984	TP-4 1-2 10-22-84 E4905	TP-5 1-2 10-22-84 E4907	TP-5 2-3 10-22-84 E4968	TP-6 1-2 10-22-84 E4909	TP-6 2-3 10-22-84 E4910	TP-7 1-2.5 10-23-84 E4916	TP-8 1-2.5 10-24-84 E4918	TP-9 1-3 10-24-64 E4920	TP-10 1-3 10-24-84 E4922	TP-11 1-3 18-24-84 E4924	TP-12 1-3 10-24-84 E4926
VOLATILE COMPOUNDS														
CHLOROBENZENE 1, 1, 1-TRICHLOROETHANE 1, 1, 2-TRICHLOROETHANE 1, 1-DICHLOROETHANE TRANS-1, 2-DICHLOROETHANE			54 <b>80</b>	360	79		11 <b>00000</b> 35000 120000				139900	9		54 <b>86</b> 55 <b>8</b> 24 <b>8</b>
ETHYLBENZENE WETHYLENE CHLORIDE TETRACHLOROETHENE TOLLENE TRICHLOROETHENE TRICHLOROETHENE	93	B 28	B 2889 2989 1680 3400	578	88	21   B 6	B 560000 140000 650000 1100000 4800000		21000 8 2900 1100 27000 6000	53 14	1500000 310000 74000 2000000 150000	76 8 15		1688 298 1288 418
VINYL CHLORIDE ACETONE 2-BUTANONE A-HETHYL-2-PENTANONE TOTAL XYLENES			50000 37000 4600	B 39000   B 33000   2500 18000		62 15 <b>9</b> 52	2000000	8900 13000 300	17 <b>000</b> 24 <b>000</b> 12 <b>000</b> 12 <b>0000</b>		650000 2800000 190000 6800000			12 <b>999</b> 12 <b>999</b>
TOTAL VOC's	192	28	187799	97330	22597	291	10505000	22450	231 <b>000</b>	67	14604000	108	130	
ACID COMPOUNDS														
PHENOL 2-METHYL PHENOL 4-METHYL PHENOL							57 <b>9099</b> 53 <b>99</b> 0							
TOTAL ACIDS	8	8	9	8	0	9	623000		8	8	8	9	•	0
BASE/NELITRAL COMPOUNDS														
1, 2-DICHLOROBENZENE ISOPHORONE NAPHTHALENE	16 <b>88</b> 27 <b>9</b>	1	1100	24 <b>00</b> 18 <b>00</b>	1798		900000 440000 180000	240	36 <b>000</b> 60000	38 <b>96</b> 47 <b>0</b> 710	78000			340
BIS(2-ETHYLHEXYL)PHTHALATE BUTYL BENZYL PHTHALATE	15 <b>000</b> 15 <b>00</b>			5788			370000	1200	61000 47000	6300 3500	59000	27 <b>900</b> 95 <b>8</b>		
DI-N-BUTYL PHTHALATE DI-N-OCTYL PHTHALATE DIMETHYL PHTHALATE	21 <b>00</b>	•	•••••	69 <b>9</b> 15 <b>00</b>	• • • • • • • • • • • •	••••••	*********		8290	340	• • • • • • • • • • • • • • • • • • • •	900	•••••	,
Fluorene Phenanthrene 2-hethylnaphthalene				450 2100							8100			
TOTAL B/N COMPOUNDS	28478	9	1100	14640	1788	9	1890000	1440	212200	15120	1451 <b>00</b>	29700	8	340
PCB's														<del></del>
AROCHLOR-1232 AROCHLOR-1268	970			340	С						39000	758	ı	
			**********	======						********	========		========	

# TABLE 4-6 SOIL ORGANIC RESULTS (ug/kg) TEST PITS SHALLOW DEPTH SAMPLES ECC Site RI Report

Sample Location: Depth (ft): Date Sampled: OTR Number:	TP-1 1-1.5 10-22-84 E4901	TP-2 1-1.5 10-22-84 E4903	TP-3 1-1.5 1 <del>8</del> -22-84 E <b>498</b> 4	TP-4 1-2 18-22-84 E4985	TP-5 1-2 19-22-84 E4987	TP-5 2-3 10-22-84 E4908	TP-6 1-2 18-22-84 E4989	TP-6 2-3 10-22-84 E4910	TP-7 1-2.5 1 <b>0</b> -23-84 E4916	TP-8 1-2.5 10-24-84 E4918	TP-9 1-3 18-24-84 E4928	TP-10 1-3 10-24-84 E4922	TP-11 1-3 18-24-84 E4924	TP-12 1-3 10-24-84 E4926
TENTATIVELY IDENTIFIED COMPOUNDS	A													
ETHYLBENZENE UNDECANE 4-METHYL-4-HYDROXYL-2-PENTANONE NONANE DECANE				20000			400000		37 <b>000</b> 7 <b>5000</b>	59 <b>00</b>				
ETHYLBENZENE ETHYL-HETHYL-BENZENE TRIDECANE PENTADECANE HEXADECANE	•••••		680	1 <b>8880</b> 2 <b>8880</b>	••••••	•••••••••••		2908	•••••	12 <b>000</b> 24 <b>000</b> 35 <b>000</b> 95 <b>00</b>	27 <b>9999</b> 27 <b>9998</b>	***********	••••••	
HEPTADECANE OCTADECANE SULFUR TOLLENE 4-METHYL-2-PENTANONE	• • • • • • • • • • • • •	•••••	•	1 <b>0000</b>	**********				•	12 <b>900</b> 47 <b>900</b> 47 <b>00</b>	14 <b>0000</b> 68 <b>0000</b>	•••••	•	4700
TETRACHLOROETHENE PHTHALATE BUTYL CELLOSOLVE t-BUTYL ALCOHOL PHENYL ETHER		•••••	• • • • • • • • • • • • • • • • • • • •		•••••			•••••	••••••••	24 <b>900</b> 9 <b>500</b>		69 12 <b>900</b>		•••••
2,6-BIS(1,1-DIMETHYLETHYL)- 2,5-CYCLOHEXADIENE-1,4-DION 2,6-BIS(1,1-DIMETHYLETHYL)- 4-METHYL PHENO 3,3,5-TRIMETHYL CYCL DHEXANONE 1,1,2-TETRACHLORDETHANE PENTANOIC ACID		•••••••••••		10000	1996	••••••	•••••	3900	••••••	•••••		12 <b>98</b> 47 <b>999</b>	•••••	
HEXANDIC ACID DIETHYL ETHER 4-HYDROXY-4-METHYL-2-PENTANONE 2-BUTANOL NOWODCOME	••••••			••••••	•••••	•••••	•••••	••••••	••••••	•••••	27 <b>8888</b>	•••••••••••••••••••••••••••••••••••••••	•••••	8 <b>00</b> 2 <b>400</b>
PHTHALIC ACID TOLLIENE-2, 4-DIJISOCYANATE 2, 4-DIJIETHYL-3-PENTANONE TETRADECANE DODECANE	19 <del>00</del> 5 <b>000</b>		688	18888	***********	•••••	500000 800000	•••••••	••••••	•••••		•••••	•••••	
1-METHYL-2-PYRROLIDINONE LAURIC ACID					************			7 <b>000</b> 1 <b>000</b>						
PERCENT MOISTURE	19.2	15.0	11.1	16.0	10.4	8.7	19.0			15.4	26,0	15. 1	13.8	10.3

- FOOTNOTES:

  A. Tentatively identified compound concentrations are estimated. A 1:1 response is assumed.

  B. Analyte has been found in the laboratory blank as well as the sample. Indicates probable contamination.

  C. Applies to pesticide parameters where the identification has been confirmed by BC/MS.

  J. Indicates an estimated value. When mass spectral data indicates the presence of a compound that meets the identification criteria and the result is less than the specified detection limit but greater than zero.

  K. Actual value, within the limitations of the method is less than the value given

#### TABLE 4-7 SOIL ORSANIC RESULTS (ug/kg) TEST PITS INTERMEDIATE DEPTH SAMPLES ECC Site RI Report

Sample Location: Depth (ft): Date Sampled: OTR Number:	TP-1 4-5 10-22-84 E4982	TP-4 2.5-3.5 10-22-84 E4906	TP-6 4-5 1 <del>8-</del> 22-84 E4911	TP-7 2.5-4 10-23-84 E4917	TP-8 2.5-4 10-24-84 E4919	TP-9 3-5 10-24-84 E4921	TP-10 3-5 10-24-84 E4923	TP-11 A 3-5 19-24-84 E4925	TP-12 3-5 1 <del>0-</del> 24-84 E4927
VOLATILE COMPOUNDS									
CHLOROBENZENE  1, 1, 1-TRICHLOROETHANE  1, 1, 2-TRICHLOROETHANE  1, 1-DICHLOROETHENE  TRANS-1, 2-DICHLOROETHENE					7790				19 <b>86</b> 62 47 9
ETHYLBENZENE NETHYLENE CHLORIDE TETRACHLOROETHENE TOLLENE	17	16	B 16	20000 4400 25000 10000	19009 1900 29008 19000	11 <b>8</b>	14 59 13 6	67	82 12 <b>9</b> 86
TRICHLORDETHENE VINYL CHLORIDE			•••••	1886	66000	13 7	ь		
ACETONE 2-BUTANONE 4-HETHYL-2-PENTANONE TOTAL XYLENES				53000 64000 100000	41908 87900 13808 41900	•			59 <b>6</b> 63 <b>8</b> 83
TOTAL VOC's	0	16	16	279286	315688	130	92	67	3649
ACID COMPOUNDS									
PHENOL 2-HETHYLPHENOL 4-HETHYLPHENOL					25000		340		
TOTAL ACIDS	0	0	8	0	25000		340	9	0
BASE/NEUTRAL COMPOUNDS									
1, 2-DICHLOROBENZENE ISOPHORONE		4488	2489	898	76 <b>888</b> 17 <b>008</b>				
I SUPPORTURE BIS (2-ETHYLHEXYL) PHTHALATE BUTYL BENZYL PHTHALATE		21 <b>98</b> 77 <b>99</b>	2680 548	640 680	12 <b>999</b> 25 <b>999</b> 59 <b>99</b>				
DI <del>-N-</del> BUTYL PHTHALATE DI <del>-N-</del> OCTYL PHTHALATE	• • • • • • • • • • • • • • • • • • • •	*******		• • • • • • • • • • • • • • • • • • • •	3900	• • • • • • • • • • • • • • • • • • • •	***********		************
DINETHYL PHTHALATE FLUORENE				2CB	1386				
PHENNYTHRENE 2-METHYLMAPHTHALENE				269 359 1900	650				
TOTAL B/N's	0	14200	5540	4720	141750	0	0	0	0
PCB¹ s									
AROCHLOR-1232 AROCHLOR-1260		548	C		1790				
TOTAL PCB's	<del></del> 0	540	9	8	1790	9	0	0	0

#### TABLE 4-7 SOIL ORGANIC RESULTS (ug/kg) TEST PITS INTERMEDIATE DEPTH SAMPLES ECC Site RI Report

Sample Location: Depth (ft): Date Sampled: OTR Number:	TP-1 4-5 10-22-84 E4982	TP-4 2.5-3.5 10-22-84 E4906	TP-6 4-5 10-22-84 E4911	TP-7 2.5-4 10-23-84 E4917	TP-8 2.5-4 10-24-84 E4919	TP-9 3-5 1 <del>8-24-84</del> E4921	TP-10 3-5 10-24-84 E4923	TP-11 A 3-5 10-24-84 E4925	TP-12 3-5 1 <b>0</b> -24-84 E4927
TENTATIVELY IDENTIFIED COMPOUNDS	A								
ethylbenzene Undecane 4 Hethyl-4—Hydroxyl-2-Pentanone Novane Decane		2 <b>0000</b> 3 <b>0000</b>		4788 5988 2486	34000 90000 45000				
ethylbenzene Ethyl-Wethyl-Benzene Tridecane Pentadecane Hexadecane	•••••••	20000	2886	12 <b>00</b> 5900 9400	11000				
neptadecane Octadecane Sulfur Toluene 4-hethyl-2-pentanone	•••••••••••••	18888		5988 6888 3588 2488	22 <b>000</b> 67 <b>900</b>		24 <b>88</b> 35 <b>000</b>		
TETROCHLORDETHENE PHTHALATE PHTHALATE LELLOSOLVE 1-BUTYL ALCOHOL PHENYL ETHER			•••••	•••••	22 <b>9999</b> 34 <b>999</b>	•	80	800	••••••
2, 5-BIS (1, 1-DIMETHYLETHYL)- 2, 5-CYCLOHEXADIENE-1, 4-DION 2, 6-BIS (1, 1-DIMETHYLETHYL)- 4-METHYLETHYLOHENC 3, 3, 5-TRIMETHYLCYCLOHEXANONE 1, 1, 2, 2-TETROCHLORGETHANE PENTANDIC ACID			••••••			••••••	248 900 2480		
HEXANDIC ACID DIETHYL ETHER 4-HYDROXY-4-WETHYL-2-PENTANONE 2-BUTANOL NONADECANE		9888	************		••••••••••	••••••	4786		•••••
PHTHALIC ACID TOLUENE-2, 4-DIISOCYANATE 2, 4-DINETHYL-3-PENTANONE TETRADECANE DODECANE	***************************************	19003	••••••	••••••	•••••••	••••••			••••••••
1-METHYL-2-PYRROLIDINONE Lauric acid				·····	· · · · · · · · · · · · · · · · · · ·	***************************************			
PERCENT MOISTURE	12.2	11.5	17.1	15.8	19.8	9. 2	15.3	16.3	18.7

### FOOTNOTES:

- A. Ientatively identified compound concentrations are estimated. A 1:1 response is assumed.

  B. Analyte has been found in the laboratory blank as well as the sample. Indicates probable contamination.

  C. Applies to pesticide parameters where the identicication has been confirmed by GC/MS.

  J. Indicates an estimated value. When mass spectral data indicates the presence of a compound that meets the identification criteria and the result is less than the specified detection limit but greater than zero.

  K. Actual value, within the limitations of the method is less than the value given

#### TABLE 4-8 SOIL BORING ORGANIC RESULTS (ug/kg) ECC Site RI Report

				INTERMEDIA	TE BORINGS					DE	EP BORINGS		7
Sample Location: Depth (ft): Date Sampled: DTR Number:	5B-01 2.5-4 10-24-84 E4912	SB-02 2.5-4 10-24-84 E4914	SB-83* 2.5-4 10-24-84 E4928	SB-03* 2.5-4 10-24-84 E4929	58-64 2-3.5 16-24-84 E4934	SB-06 2-3.5 10-24-84 E4932	58-88 2.5-4 18-24-84 E4931	SB-09 2.5-4 10-24-84 EB077	SB0104 5.5-7 10-24-84 E4913	SB8284 5.5-7 18-24-84 E4915	SB0403 5-6.5 10-24-84 E4933	SB8885 7-8-5 19-24-84 E4935	SB8984 5.7-7 18-24-84 E4938
VOLATILE COMPOUNDS								<del></del>					
1, 1, 1-TRICHLOROETHANE 1, 1-DICHLOROETHANE 1, 1, 2-TRICHLOROETHANE CHLOROFON 1, 1-DICHLOROETHENE	14 57	49000 2900 1600	11 <b>900</b> 150	65	3 J 14	27 <b>000</b>	27 J	18 <b>800</b> 380 J				11 5 J	
TRANS-1, 2-DICHLOROETHENE ETHYLBENZENE NETHYLENE CHLORIDE TETRACHLOROETHENE TOLUENE	37 15 1 <b>99</b> 44 52	1508 21900 8 19000 E 11908 31900	3 19 <b>00</b> 68 <b>0</b>	74	17 8 5 J	4800 4100 16000 11000	72 27 J 59 B 26 J 178	1858 2888	27 B	34 B 1 <b>0</b>	33	41 54 8 14	29 190 120
Trichloroethene Acetone 2-Butanone 2-Hexanone	39 1488 1298	68 <b>889</b> 17 <b>99</b> 9	348 32998 24 <b>88</b> 9	558 550	16 6 J 78	119000 17000 8000 JE	1600	18888 6688 B 928	66		18 B	3 J 41 E	3 6500 1000
4-HETHYL-2-PENTANONE TOTAL XYLENES	250 95	118000		36	36	21888	35 J 190	2888				11	44
TOTAL VOC's	3303	12900	70979	1275	175	220900	3912	68390	27	34	51	188	8069
ACID COMPOUNDS													
PHENOL 2-METHYLPHENOL 4-METHYLPHENOL	-					61 <b>8</b>		1198					
TOTAL ACIDS	0	0	8	8	8	619	8	1106	8	9	8	8	8
BASE/NEUTRAL COMPOUNDS													
ISDPHORONE NAPHTHALENE NISY(2-ETHYLHEXYL) PHTHALATE BISY(2-ETHYLHEXYL) PHTHALATE BUTYL ENTYL PHTHALATE DI-N-BUTYL PHTHALATE	239	640			42 <b>0</b> JI	<b>5848</b> В <b>4848</b> Л	73 <b>0</b> 8 53	498 J 329 JB			31 <b>0</b> E	270 .	I
DIETHYL PHTHALATE DIMETHYL PHTHALATE	••••••	9000 1200	• • • • • • • • • • • • • • • • • • • •	••••••	•••••	12 <b>99</b> 3 <b>69</b> J		•••••	**********	•••••	***********	•••••	
TOTAL B/N COMPOUNDS	239	10840	8	9	420	2468	783	720	9	9	9	0	<del></del>

## TABLE 4-8 SOIL BORING ORGANIC RESULTS (ug/kg) ECC Site RI Report

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				INTERNEDIA	TE BORINGS				DEEP BORINGS				•
Sample Location: Depth (ft): Date Sampled: OTR Number:	SB-01 2.5-4 10-24-84 E4912	SB-82 2.5-4 18-24-84 E4914	SB-83* 2.5-4 18-24-84 E4928	58-03# 2.5-4 18-24-84 E4929	58-84 2-3.5 10-24-84 E4934	SB-86 2-3.5 10-24-84 E4932	58-88 2.5-4 18-24-84 E4931	SB-09 2.5-4 10-24-84 EB077	580104 5.5-7 10-24-84 E4913	SB6264 5.5-7 18-24-84 E4915	SB8483 5-6.5 18-24-64 E4933	SB0805 7-8.5 10-24-84 E4935	SB0904 5.7-7 10-24-84 E4930
PESTICIDE COMPOUNDS													
NONE DETECTED						*********			. <del></del>			***	
PCB¹ s													
NONE DETECTED								######################################	• •				
TENTATIVELY IDENTIFIED COMPOUNDS													
DECANE UNDECONE TRICHLOROFLUOROMETHANE 4-METHYL-2-PENTANOL TETRACHLOROETHENE		900 1000			<b>4</b> :	j J					10 .	J 12 .	ı
1, 1, 2-TRICHLORO- 1, 2, 2-TRIFLUOROETHANE ISOPROPYL ALCOHOL 2-BUTANOL DIETHYL ETHER HEXANE	•••••	• • • • • • • • • • • • •	•••••	119 9 <b>9</b>	1		•••••	2 <b>4000</b> J	• ••••••	•••••	••••••	•	48 J 58
PERCENT MOISTURE	13.7	11.4	11.59	11.86	12	19	12	8	10.7	<del></del>	11	8	14.5

#### FOOTNOTES:

\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

- ES:
  A. Tentatively identified compound concentrations are estimated. A 1:1 response is assumed.
  B. Analyte has been found in the laboratory blank as well as the sample. Indicates probable contamination.
  C. Applies to pesticide parameters where the identicication has been confirmed by GC/MS.
  J. Indicates an estimated value. When mass spectral data indicates the presence of a compound that meets the identification criteria and the result is less than the specified detection limit but greater than zero.
  K. Actual value, within the limitations of the method is less than the value given
  Duplicate samples were taken at SB-83

The specific compounds detected, their maximum reported concentration, and general occurrence onsite are summarized in Table 4-9. Site cleanup activities completed after the Phase 1 sampling included soil removal from the northern drum storage area, capping of the site with berm soil, and general onsite soil disturbance associated with waste handling and removal. Surface cleanup activities have therefore removed and/or redistributed some portion of the soil sampled during the Phase 1 sampling. The information collected for analysis of Phase 1 samples is presented to indicate the types, concentrations, and general site location of organic contaminants once present in soil at the site.

Organic Contamination, Phase 2 Sampling. As with the Phase 1 samples, analysis of soil samples collected during Phase 2 sampling activities detected a wide variety of organic contaminants. Major compound groups detected included volatile organics, phenols, phthalates, polynuclear aromatic hydrcarbons (PAHs), and PCBs. Of these compound groups, volatile organics and phthalates were more commonly detected and generally were reported at the highest concentrations. Figures 4-5, 4-6, and 4-7 summarize the distribution of the major organic compound groups detected in Phase 2 soil samples.

Table 4-9
SUMMARY OF ORGANIC COMPOUNDS DETECTED IN PHASE 1 SOIL SAMPLES

		Sit	e Areas Where Compoud was Detecte	d in Phase 1 Samples
			Drum and Tank	Soil Areas South
			Storage Area North	of the Cooling Water
	Maximum Observed	Berm	of the Cooling	Pond and Adjacent
	Concentration (ug/kg)	Area	Water Pond	to the Concrete Pond
Volatiles				
1,2-Dichlorethane	280			X
1,1,-Trichlorethane	7,411,400		X	X
1,1-Dichloroethane	700			X
Chloroform	41,800		Х	X
Trans-1,2-Dichloroethene	79,700		X	X
Cis-1,2-Dichlorepropane	12,000			х
Ethylbenzene	5,649,000		х	X
Methylene Chloride	515,000	X	X	X
Chloromethane	70	X		
Tetrachloroethene	4,116,000		Х	Х
Toluene	954,000		Х	X
Trichloroethene	6,080,200		X	X
Vinyl Chloride	6,400			X
Acetone	30,300			x
2-Butanone	99,200		Х	Х
4-mety1-2-Pentanone	29,600		Х	X
Styrene	19,000		х	X
Total Xylenes	1,160,000		X	х
Acid Extractable Compounds				
2,2-Diemthyl phenol	88,000			x
Phenol	447,000		X	X
Benzoic Acid	28,200		X	X
2-Methyl Phenol	142,600		X	х
4-Methyl Phenol	535,600		x	x

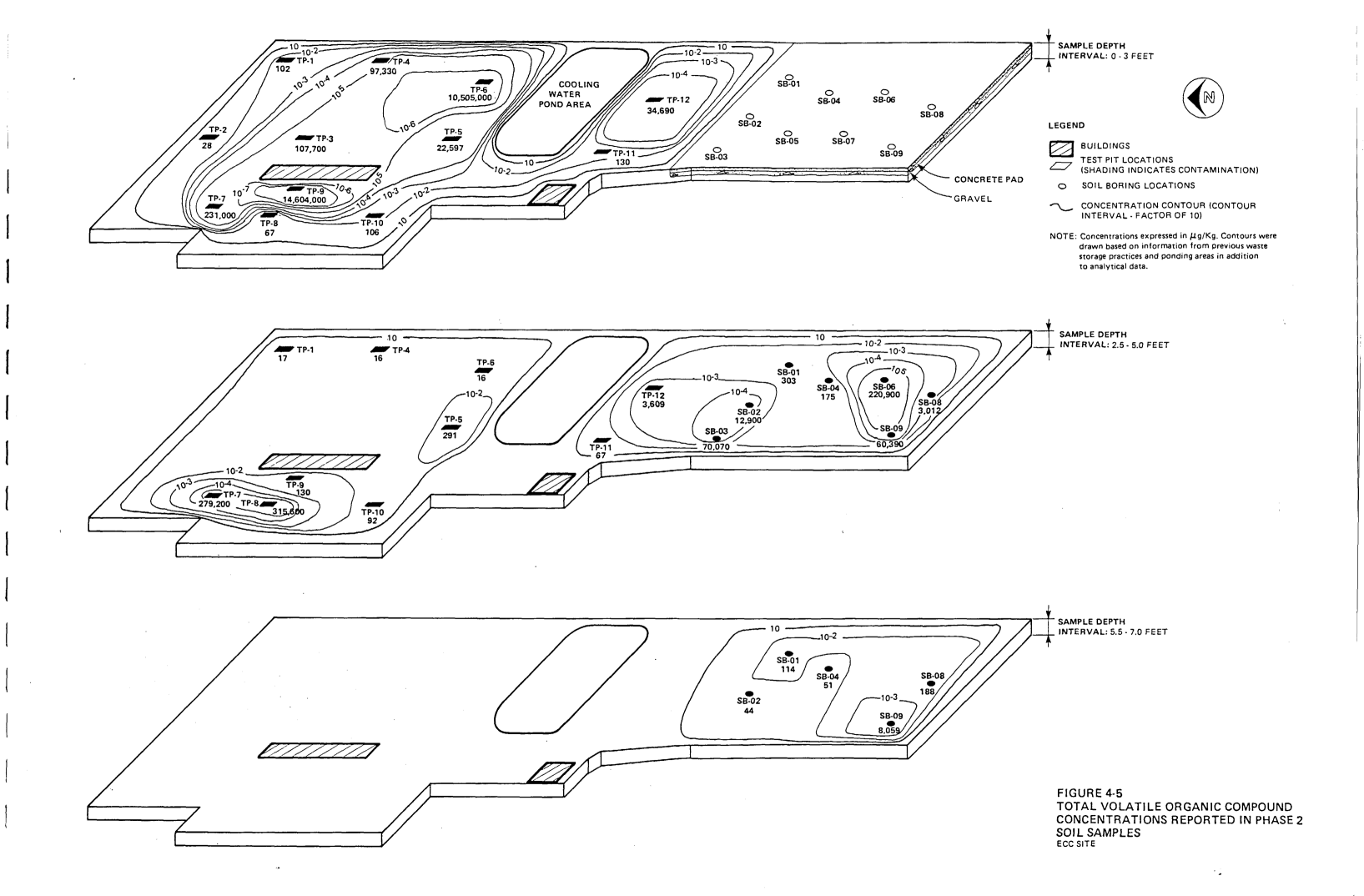
## Table 4-9 (Continued)

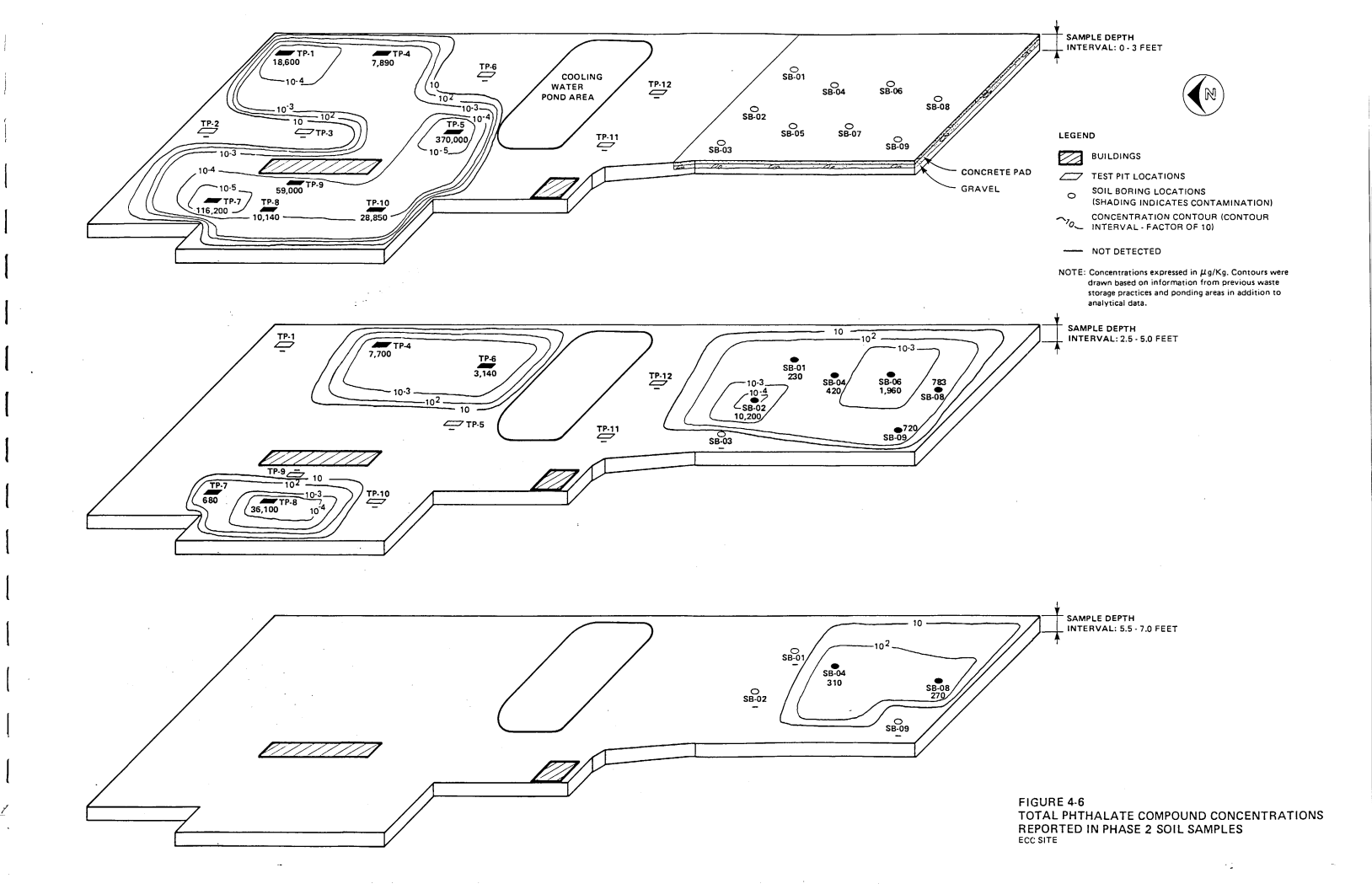
		Sit	e Areas Where Compoud was Detecte	ed in Phase 1 Samples
		<del></del>	Drum and Tank	Soil Areas South
			Storage Area North	of the Cooling Water
	Maximum Observed	Berm	of the Cooling	Pond and Adjacent
	Concentration (ug/kg)	Area	Water Pond	to the Concrete Pond
Base/Neutral Extractable				
1,2,4-Trichlorobenzene	389,600			
1,2-Dichlorobenze	2,160,000		X	X
1,4-Dichloroe benzene	570,000			X
1,2-Diphenylhydrazine	68,600K		X	X
Hexachloroebutadiene	5,000			X
Isophorone	409,200		X	Х
Naphthalene	470,000		X	X
Nitrobenzene	7,800			Х
N-Nitrosodimethyamine	9,900			Х
N-Nitrosodiphenylamine	1,400	X		
N-Nitrosodipropylamine	12,000			X
Bis(2-ethylhexyl)phthalate	3,800,000	Х	X	Х
Benzyl Butyl Phthalate	1,282,000		X	Х
Di-N-Butyl Phthalate	112,200		X	X
Di-N-Octyl Phthalate	300,000		x	X
Dicthyl Phthalate	3,500			X
Dimethyl Phthalate	25,400		х	
Phenanthrene Phthalate	8,000		X	
2-Methylnapthalene	130,000		X	Х
Pesticides				
Delta-BHC	760		Х	х
Gamma-BHC (lindane)	170		X	
Heptachlor	210		X	
Aldrin	20		X	
Dieldrin	700	X	X	X
Endrin	11,200		X	x
Endosulfan I	8,300			X
Endosulfan II	11,100		x	х
4,4-DDD	5,900		X	
4,4-DDE	830		x	X
4,4-DDT	36,000	X	X	Х

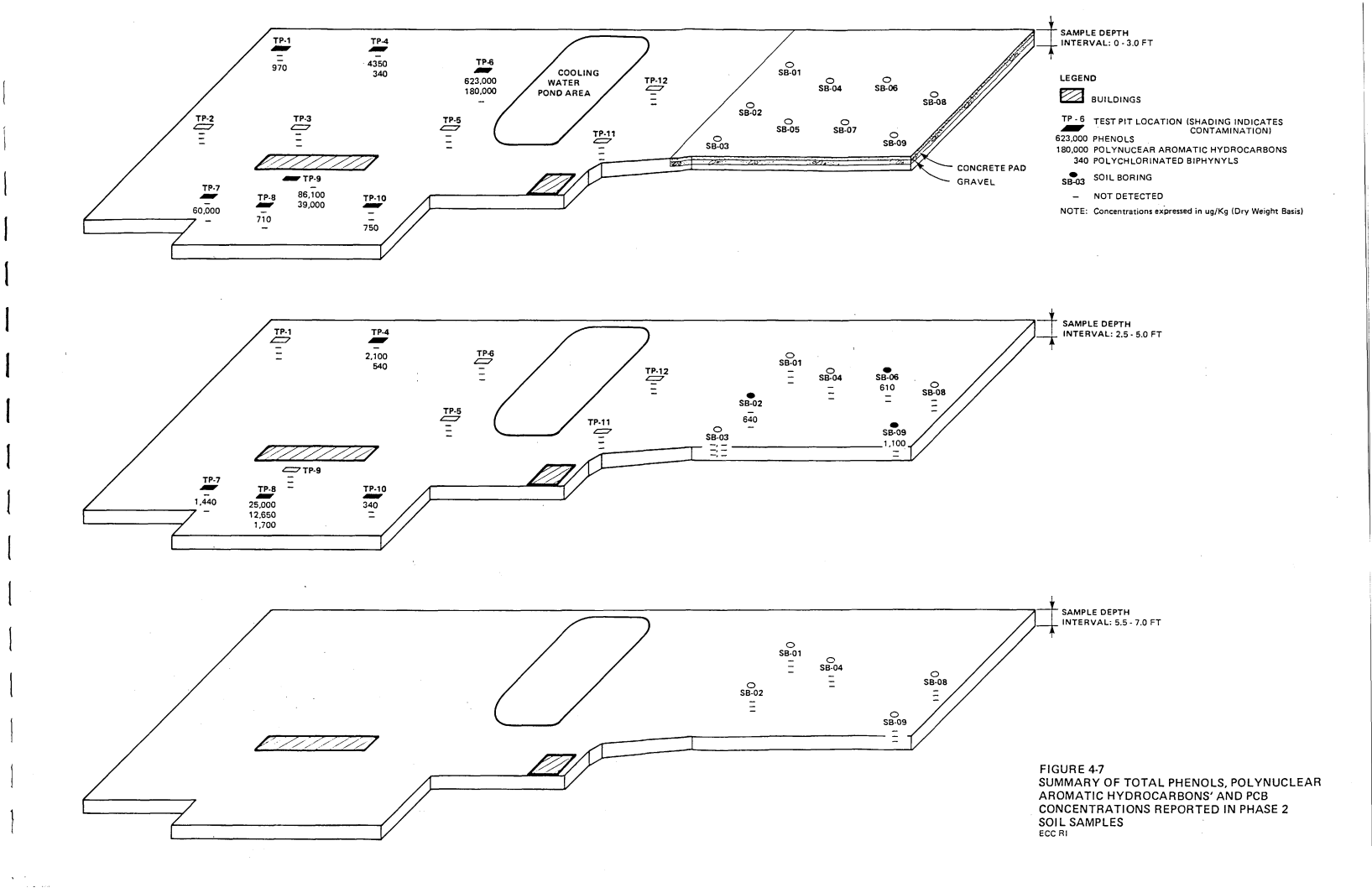
Table 4-9 (Continued)

		Site Areas Where Compoud was Detected in Phase 1 Samples		
	Maximum Observed Concentration (ug/kg)	Berm Area	Drum and Tank Storage Area North of the Cooling Water Pond	Soil Areas South of the Cooling Water Pond and Adjacent to the Concrete Pond
Pesticides (Continued)				
Endrin Aldehyde	20,000		X	X
Endosulfan Sulfate	19,000		x	x
Chlordane	2,700			X
Toxaphene	10,800			X
PCB's				
Arochlor-1016	10,800			X
Arochlor-1232	16,200			X
Arochlor-1248	10,800			Х

GLT360/72







Nineteen VOCs were detected in soil samples from the site. The primary VOC's detected in soil samples from the site include the following:

1,1,1-Trichloroethane

Methylene Chloride

Tetrachloroethene

Acetone

Trichloroethene

2-Butanone

Ethylbenzene

4-methyl-2-Pentanone

Toluene

Xylenes

Volatile organic compounds are the most widespread organic contaminant at the site and were detected to the maximum soil sampling depth of 8.5 feet. Except for areas near test pits 7 and 8 and below the concrete pad, total VOC concentration in subsurface soil (2.5-8.5 feet) are generally several orders-of-magnitude lower than observed in surface soil.

Phthalate compounds detected in soil samples at the site are:

Bis (2-ethylhexyl) phthalate Di-n-octyl phthalate

Butyl Benzyl Phthalate

Diethyl Phthalate

Di-n-butyl Phthalate

Dimethyl Phthalate

The distribution of phthalate compounds is similar to that of the VOC's, except that phthalates are generally reported in lower concentrations and are not as frequently detected in subsurface soils. As with the VOC's, phthalate compound concentrations in subsurface soil are generally several orders-of-magnitude less than detected in surface soil.

Acid extractable compounds detected in soil from the site are:

2,4-Dimethyphenol

Phenol

2-Methylphenol

Benzoic Acid

4-Methylphenol

Phenol was the most frequently detected of these compounds. Contamination of soil with these compounds appears to be limited to localized areas; surface soil in the vicinity of test pit 6; surface soil adjacent to the concrete pad; subsurface soil in the vicinity of test pit 8; and subsurface soil beneath the concrete pad.

PAH's detected in soil at the site are:

Napthalene

Fluorene

Phenanthrene

3-Methylnaphthalene

Naphthalene is the most frequently detected PAH and the only PAH detected in soil samples from beneath the concrete pad. The detection of PAH compounds is, except for one sample, limited to surface soil adjacent to the concrete pad and soil in the northern drum and tank storage areas.

PCB's were detected in only six Phase 2 soil samples. Their detection was limited to soil sampled in the northern drum and tank storage areas. The maximum concentration reported was 39,000 ug/kg, but concentrations were generally less than 1,000 ug/kg.

### CONCLUSIONS AND OBSERVATIONS

Inorganic contamination of the soil is apparently greatest in the near surface (0-3 feet) soil in northern portions of the site. Inorganic contamination does appear to extend to depths of at least 5 feet in the northern portions of the site, although it is less widespread than observed in the overlying shallow soil.

General observations regarding the organic contamination at the site are:

o Primary organic contaminants at the site are VOC's and phthalates. These compound groups are the

most widespread organic contaminants and are generally present in the highest concentrations.

Organic contamination decreases in the variety of compounds and their associated concentrations with depth. However, organic contaminants were detected to the maximum depth of sample analysis (8.5 feet).

## HYDROGEOLOGIC INVESTIGATIONS

Boone County, Indiana, is in a physiographic unit known as the Tipton Till Plain, a nearly flat to gently rolling glacial plain, which is the result of continental ice sheets that covered the county about 20,000 years ago. During the period, known as the Pleistocene Epoch, large quantities of earth materials were deposited upon the bedrock surface, with a maximum thickness approaching 350 feet. The major aquifers in Boone County are in sand and gravel deposits of glacial origin. These deposits are also important sources of aggregate materials.

The bedrock formations beneath the glacial drift in Boone County consist of limestones and dolomites of Silurian and Devonian age and shales of Devonian and Mississippian age. The beds generally dip about 10 to 30 feet per mile to the southwest toward the Illinois Basin.

## SCOPE AND METHODS

A hydrogeologic investigation was conducted to define the soil stratigraphy, characterize aquifer conditions and determine groundwater flow directions, gradients, seasonal water level variations in the vicinity of the ECC site, and to define subsurface contaminant migration and pathways.

The program included an electrical resistivity survey, test

drilling with soil sampling and rock coring, installation of monitoring wells and sampling of groundwater. Details on methods and results are presented in TM 3-1 and 3-2 of Appendix A.

## Electrical Resistivity Survey

An electrical resistivity survey was conducted to investigate the presence and lateral continuity of shallow sand and gravel deposits and the presence of fine-grained glacial tills in the vicinity of the ECC site. A secondary objective was to investigate the presence of a groundwater contaminant plume. Due to the presence of many surface features that may interfere with electrical resistivity, vertical electrical soundings were taken at each of 52 stations surrounding the site.

## Test Drilling

A series of monitoring well clusters were installed around the ECC site using hollow stem augers and/or rotary techniques. The wells were classified into three groups based on their relative borehole depths. Eleven shallow boreholes (wells) were drilled to a maximum depth of about 30 feet (designated "A"). One intermediate borehole (well) was drilled to approximately 100 feet (designated "B"). Four deep boreholes (wells) were drilled into the top of rock,

approximately 155 to 165 feet (designated "C"). Borehole locations are shown in Figure 4-8. Continuous split-spoon samples were taken at 2 foot intervals in the upper 20 to 30 feet in one borehole at each cluster and at 5 foot intervals thereafter to top of rock. The boreholes were drilled in three phases. Phase 1 included boreholes 1A, 1C, 2A, 2B, 2C, 3A, 3C, 4A, 4C, and 5A. Phase 2 included 6A and 7A which replaced 4A due to a drilling contamination problem. Phase 3 included 8A, 9A, 10A, and 11A.

Laboratory testing included index tests for soil identification and classification. These consisted of Atterberg limits, moisture contents and mechanical grain size analysis. Samples were selected for testing after visual classification of all samples from a borehole and were selected on the basis of being representative of soil types encountered.

## Monitoring Well Installation

A total of 16 2-inch diameter PVC monitoring wells were. installed in the boreholes discussed above. Wells were developed either by flushing with clean water or by air lifting. The deep and intermediate wells (1C, 2B, 2C, 3C, and 4C) were free flowing and a packer assembly was devised to control the well water flow.



LEGEND

REMEDIAL INVESTIGATION MONITORING WELL ECC-7A

NOTE: All well locations are approximate

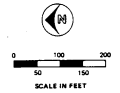


FIGURE 4-8
MONITORING WELL LOCATIONS
ECC RI REPORT

## Monitoring Well Sampling

Monitoring wells were sampled in three phases. Phase 1, (July 18 and 19, 1983) included sampling of 1A, 1C, 2A, 2B, 2C, 3A, 3C, 4C, and 5A. Phase 2 (November 29 and 30, 1983) sampling included wells 1A, 1C, 2A, 2B, 2C, 3A, 3C, 5A, 6A, and 7A. Phase 3 (December 12 and 13, 1984) sampling included 1A, 2A, 3A, 5A, 6A, 7A, 8A, 9A, 10A, and 11A. The deep and intermediate wells were purged and sampled by opening the check valve in the packer assembly. The shallow wells were purged and sampled with a submersible stainless steel pump. Samples for VOC analysis were obtained with a stainless steel bailer. At least three well volumes were purged from each well prior to sampling. Samples for inorganic analysis were filtered in the field through a 0.45 micron filter and then preserved with nitric acid.

Water levels were taken using an electric well sounder. In the flowing deep and intermediate wells, 1-1/4 inch PVC pipe extensions were added to the packer assembly until the potentiometric surface was obtained.

## Residential Well Sampling

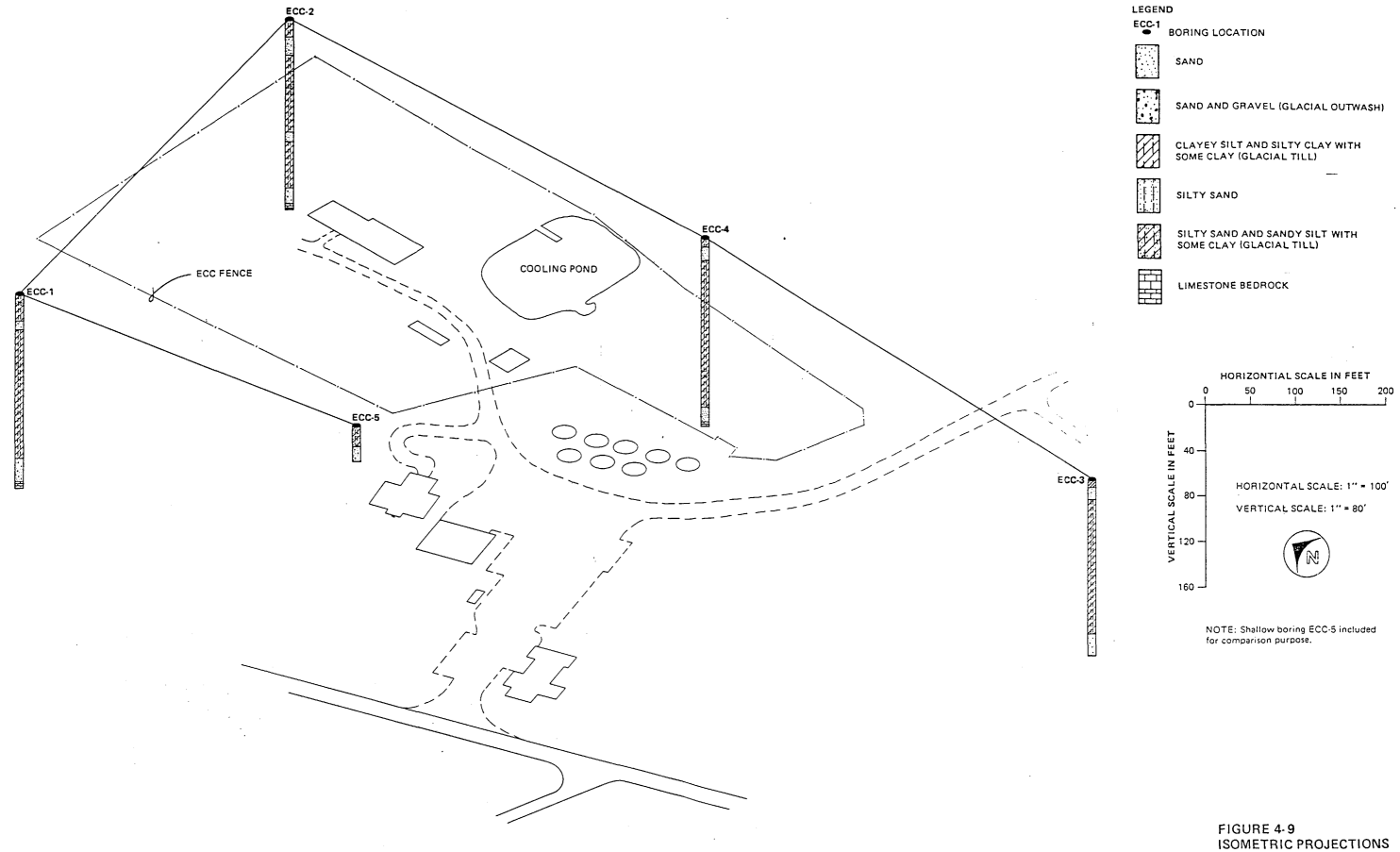
Five residential wells were sampled on May 10, 1983. Wells were pumped for 20 to 30 minutes prior to sampling. Samples were collected by filling the bottles directly from the

faucet closest to the well head. Inorganic samples were not field filtered prior to preservation with nitric acid.

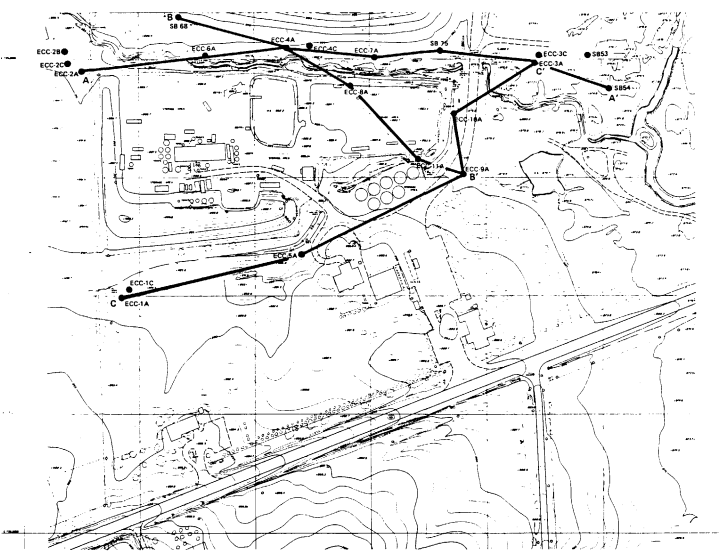
## RESULTS

## Site Geology

Soil types encountered at ECC from the ground surface to the top of rock consist of glacial tills, glacial outwash and possibly some shallow alluvial deposits. Figure 4-9 illustrates soil types for the four deep borings. The glacial till deposits, consisting predominantly of clayey silt and silty clay, formed the thickest sequence encountered. They appear to be highly overconsolidated based on Atterberg limits and relatively low permeability. Sands and gravels were found at nearly all boring locations. These consist of fine to coarse sand and gravel that are highly permeable. Some alluvial deposits occur near the ground surface, especially near the southeast corner of the ECC site and generally consist of fine sand and silty sand. Cross sections were prepared illustrating shallow soil conditions at the site (see Figure 4-10). Cross sections are presented in Figures 4-11, 4-12, and 4-13. Included are some of the borings completed previously at NSL. The shallow soil stratigraphy appears to be very complex near the south end of the ECC site. This is probably due to the combination of till, outwash and alluvial deposits present in this area.



OF DEEP BORINGS ECC RI REPORT



LEGEND

REMEDIAL INVESTIGATION MONITORING WELL ECC-7A

A \_\_\_\_A'CROSS SECTION LOCATION

NOTE: All well locations are approximate

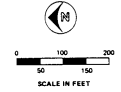
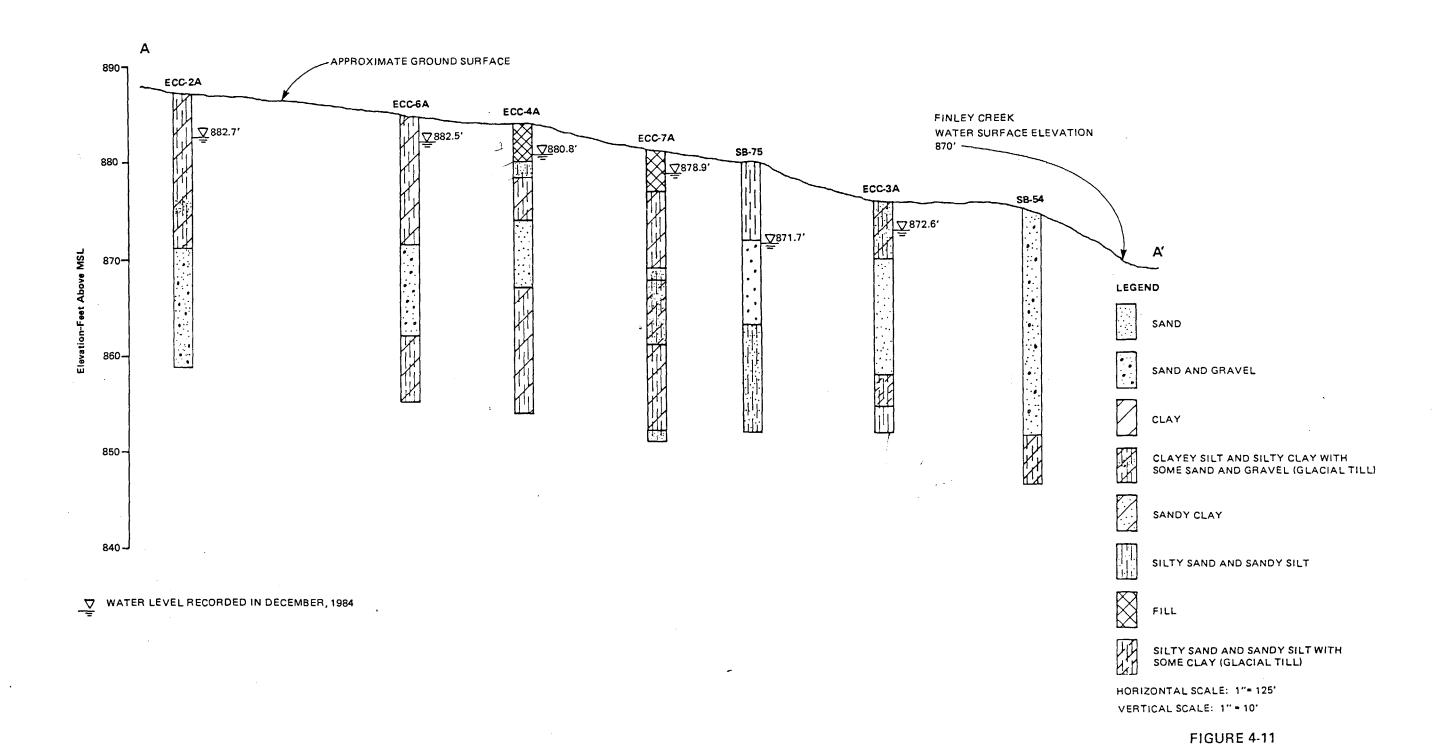
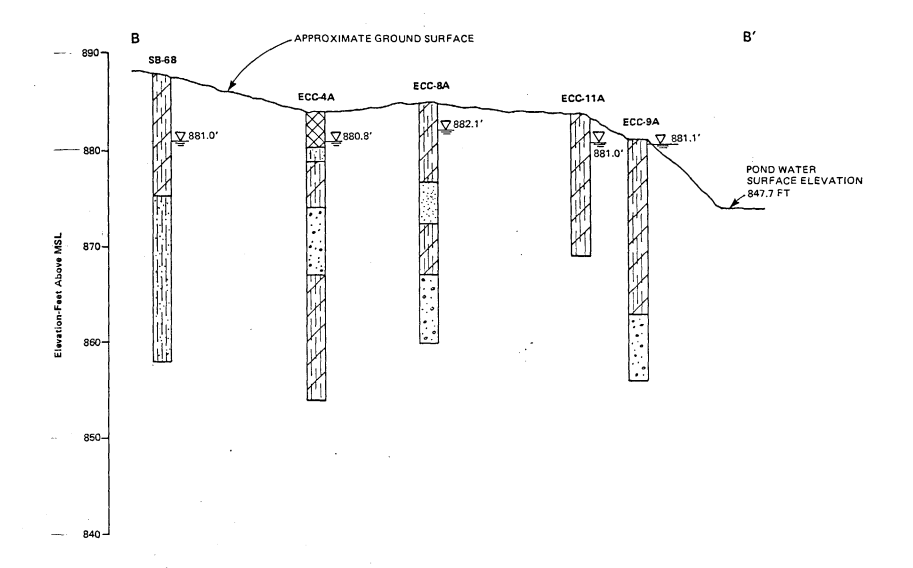


FIGURE 4-10 SOIL BORING CROSS SECTION LOCATIONS ECC RI REPORT



CROSS SECTION A-A'

ALONG THE UNNAMED DITCH ECC RI REPORT



▼ WATER LEVEL RECORDED IN DECEMBER, 1984

### LEGEND

SAND



SAND AND GRAVEL



CLAY



CLAYEY SILT AND SILTY CLAY WITH SOME SAND AND GRAVEL (GLACIAL TILL)



SANDY CLAY



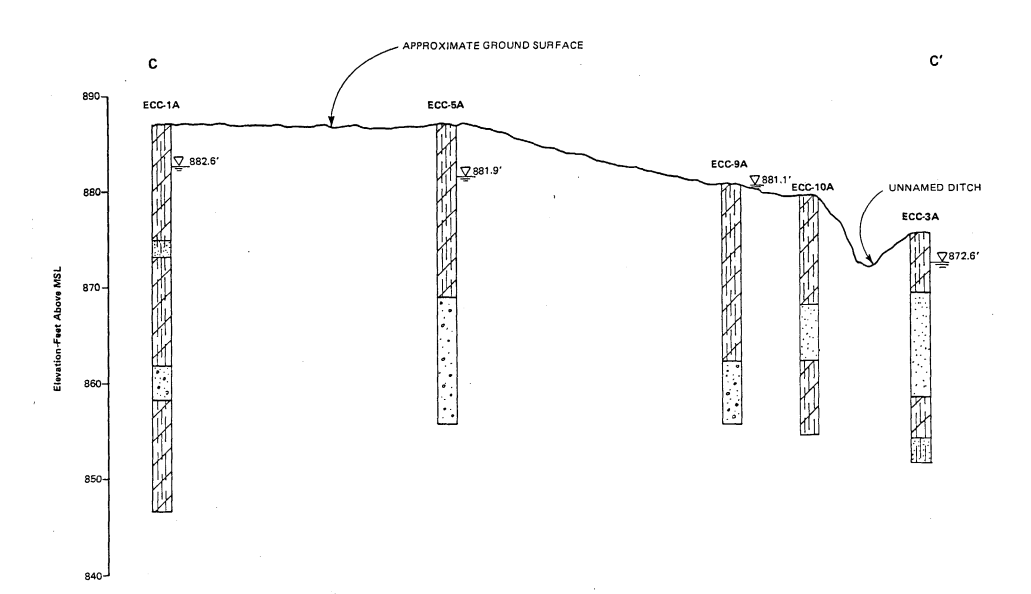
SILTY SAND AND SANDY SILT



FILL

HORIZONTAL SCALE: 1"= 125' VERTICAL SCALE: 1" = 10'

> FIGURE 4-12 CROSS SECTION B-B' NORTHEAST TO SOUTHWEST ACROSS SITE ECC RI REPORT



SAND

SAND AND GRAVEL

CLAY

CLAYEY SILT AND SILTY CLAY WITH SOME SAND AND GRAVEL (GLACIAL TILL)

SANDY CLAY

SILTY SAND AND SANDY SILT

FILL

HORIZONTAL SCALE: 1"= 125'

VERTICAL SCALE: 1"= 10'

LEGEND

▼ WATER LEVEL RECORDED IN DECEMBER, 1984

FIGURE 4-13
CROSS SECTION C-C'
NORTHWEST TO SOUTHEAST
ACROSS SITE
ECC RI REPORT

# Hydrogeologic Units

Four hydrogeologic units occur at different elevations beneath the site. These are:

- o A Shallow saturated zone consisting of clayey silts and silty clays approximately 5 to 15 feet below ground surface. The lithology of this unit is areally heterogeneous.
- o A sand and gravel zone, approximately 20 to 30 feet below ground surface, that may be semiconfined in places.
- o A thick zone of clayey silts and silty clays, approximately 30 to 150 ft below ground surface. This unit appears to act as an aquitard.
- o A deep confined aquifer consisting of sand and gravel, approximately 150 to 165 feet below ground surface

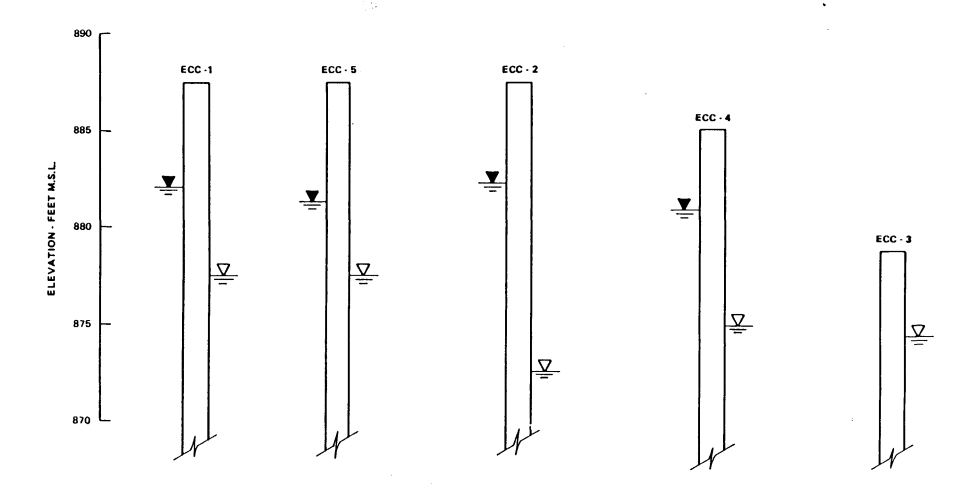
Shallow Saturated Zone. The approximate depth to the water table was identified while drilling with hollow-stem augers and continuous split-spoon sampling. Depths to the water table ranged from 6 feet at ECC-3 to approximately 10 feet at ECC-1, 4 and 5, and to 15 feet at ECC-2. The water table

occurred in fine-grained soils, usually clayey silts or silty clays at ECC-1, 2, 4, 5, 6 and 7. At ECC-3, it occurred in a fine sand, relatively free of silt.

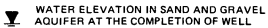
The majority of shallow wells are completed in the sand and gravel zone below the uppermost hydrologic unit. Therefore, water levels in these wells may not represent the water table. In addition, the approximate depth to the water table was identified during drilling of these test borings. The difference in elevation is shown in Figure 4-14.

Monitoring wells 3A and 11A are completed in the uppermost hydrogeologic unit and the water level data collected from these wells represents the water table.

Sand and Gravel Aquifer. An areally extensive sand and gravel zone was identified between approximately the 20- and 30-foot depth at ECC-1, 2, 4, 5, 6, 8, 9, and 10. The potentiometric surface of this zone is at a higher elevation than the water table at these boring locations. This zone appears to be a glacial outwash sand and gravel zone, overlain by a silty clay till which, in places, may act as an aquitard. The upper till unit appears to be 10 to 15 feet thick throughout the northern half of the ECC site. At ECC-3, the shallow sand and gravel aquifer was overlain by 5 feet of till. The potentiometric surface of the sand and gravel zone at this well was not found to be appreciably different during drilling of the test boring. The shallow



#### LEGEND



NOTE: Shallow sand and gravel aquifer was not encounteded at ECC - 3

**VERTICAL SCALE 1" = 5'** 

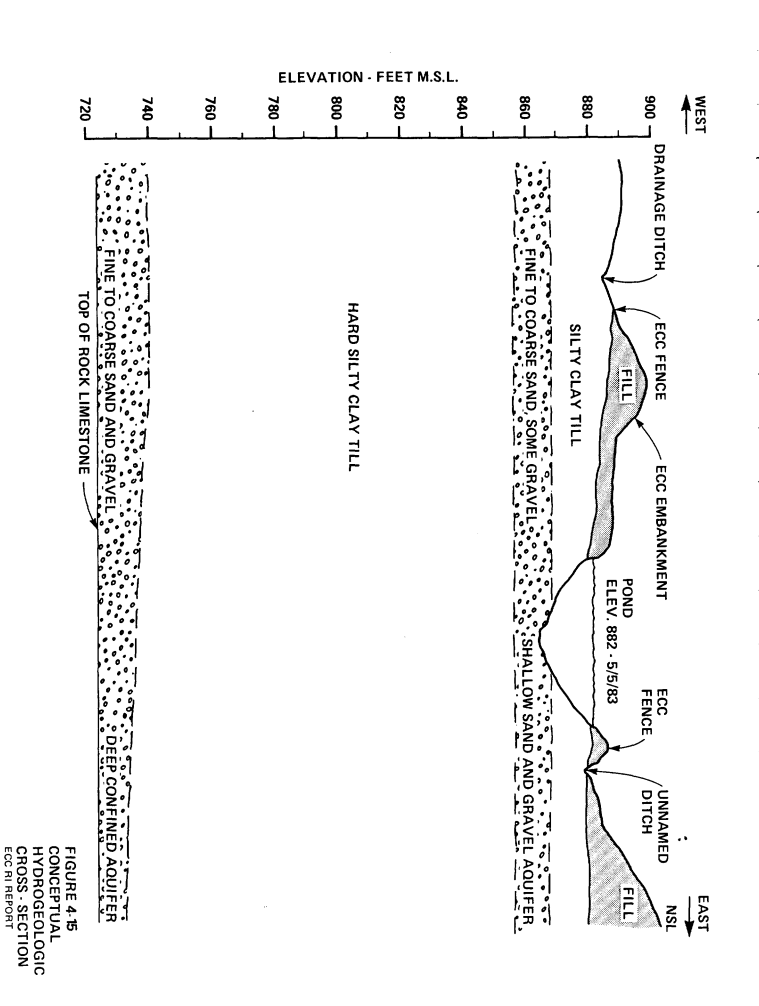
**HORIZONTAL - NOT TO SCALE** 

FIGURE 4-14
HEAD DIFFERENCE BETWEEN THE SHALLOW
SATURATED ZONE AND THE SAND AND
GRAVEL AQUIFER
ECC RI REPORT

sand and gravel zone at ECC-4 occurs at a higher elevation than at ECC-1, 2, and 5, and the zone consists of a finer, silty sand at ECC-4 than at the other boring locations. The shallow sand and gravel zone identified at the ECC-6, ECC-8, and ECC-9 locations has very similar characteristics to the 20- to 30-foot depth at ECC-1, 2, and 5. At ECC-7, the zone is similar to ECC-4, with large amounts of silt and interbedded clay lenses.

The cooling water pond appears to be excavated below the top of the shallow sand and gravel aquifer as shown in Figure 4-15. Groundwater inflows to the cooling pond were reported to be about 2,500 gallons/hr during the dewatering operation performed by the surface cleanup contractor. This high influx indicates that pond was excavated into the top of the shallow sand and gravel aquifer.

Deep Confined Aquifer. A deep confined zone was found in outwash sands and gravels near the top of rock in all four deep borings (see Figure 4-9). The potentiometric surface of this zone is above ground surface throughout the site This aquifer is confined by an extensive sequence of overlying till, which consists of very stiff to hard clayey silts and silty clays with very low permeabilities (based upon Atterberg limits and visual classification). The natural moisture contents and Atterberg limits indicate that this till is highly overconsolidated.



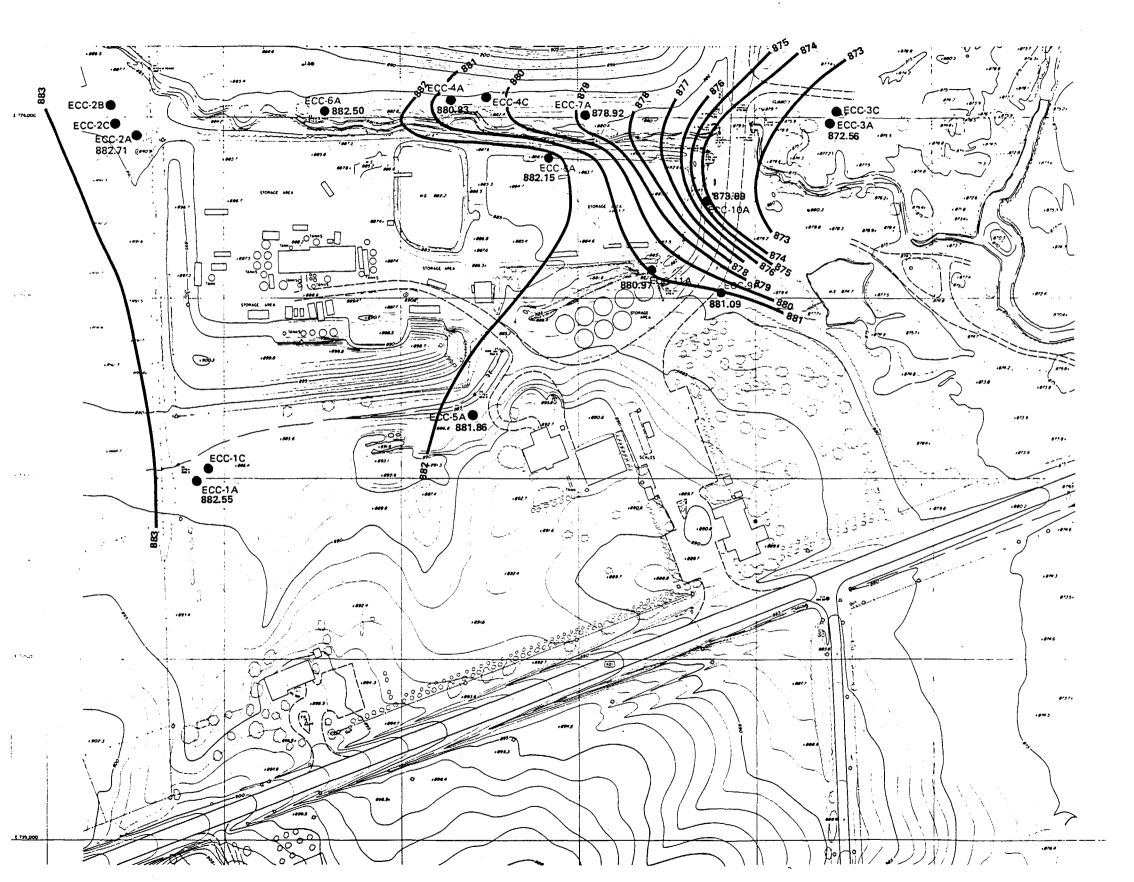
Other Hydrogeologic Units. Several discontinuous sandy zones occur in the till and are water-bearing zones.

Monitoring well ECC-2B is completed in such a zone, approximately 100 feet below ground surface. The water level in ECC-2B is very close to the water level in the deep well, ECC-2C. This zone is about 10 feet thick; however, other zones encountered were usually less than 5 feet thick and generally contained considerable amounts of silt and clay.

# Groundwater Flow System

Interpretation of the shallow groundwater flow system at the site is difficult because of the heterogeneity of the geologic materials and because of the man-induced changes to the local hydrologic system.

Although the geologic materials of the upper two hydrogeologic units are dissimilar, they appear to be hydraulically connected. Interpretation of the shallow groundwater flow system is shown in Figure 4-16. Table 4-10 presents groundwater elevations for ECC wells. Groundwater below the site generally appears to travel south and discharge into Finley Creek or the unnamed ditch near its confluence with Finley Creek. Along the eastern edge of the southern half of the site groundwater appears to flow in an eastern direction and discharge into the unnamed ditch.



# LEGEND

● REMEDIAL INVESTIGATION MONITORING WELL ECC-7A

=882 = CONTOURS FOR DECEMBER 1984 DATA

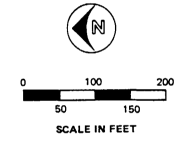


FIGURE 4-16
GROUNDWATER CONTOUR MAP
DECEMBER 1984
ECC RI REPORT

# Table 4-10 (Page 1 of 2) GROUNDWATER LEVELS IN RI MONITORING WELLS ECC SITE

• *	Ground Surface	Top Casing	Feet from	Flowation	Data
Well No.	Elevation Ft MSL	Elevation Ft MSL	Ground a Surface	Elevation Ft MSL	Date Recorded
WEIL NO.	rc. non	<u></u>	<u> </u>	<u></u>	
ECC-1A	887.13	890.13	-5.46	881.67	6/29/83
			-5.67	881.46	7/19/83
			-6.24	880.89	9/1/83
			-5.45	881.68	11/29/83
			-4.58	882.55	12/12/84
ECC-1C	886.76	889.46	+5.06	891.82	6/29/83
			+4.70	891.46	7/18/83
			+3.99	890.75	11/29/83
			+2.50	889.26	12/13/84
ECC-2A	887.21	890.21	-5.15	882.06	6/29/83
DOC 241	00,022	373422	-5.43	881.78	7/19/83
			-6.15	881.06	9/1/83
			-5.31	881.90	11/29/83
			-4.50	882.71	12/12/84
ECC-2B	886.65	889.65	+5.19	891.84	6/29/83
DCC 22	00000	****	+4.34	890.99	7/20/83
			+3.78	890.43	11/29/83
			+2.10	888.75	12/13/84
ECC-2C	886.80	889.70	+5.09	891.89	6/29/83
100 20	000.00	00,1,0	+4.78	891.58	7/18/83
			+3.78	890.67	11/29/83
			+2.29	889.09	12/13/84
ECC-3A	876.47	878.87	-4.31	872.16	6/29/83
DCC 3A	07011	3,333,	-5.13	871.34	7/19/83
			<b>-4.</b> 90	871.57	9/1/83
			-5.26	871.21	11/29/83
			-3.91	872.56	12/12/84
ECC-3C	877.19	879.59	+12.52	889.71	6/29/83
LCC 3C	07,100		+12.24	889.43	7/20/83
			+13.30	890.49	11/30/83
ECC-4A	884.34	887.24	<b>-4.</b> 11	880.23	6/29/83
ECC 4A	004,0		-4.38	879.96	7/19/83
			-4.66	879.68	9/1/83
			-3.51	880.83	12/12/84

Table 4-10 (Page 2 of 2)

Welf'No.	Ground Surface Elevation Ft MSL	Top Casing Elevation Ft MSL	Feet from Ground Surface	Elevation Ft MSL	Date Recorded
ECC-4C	884.54	887.24	+7.71 +6.93 +6.10 +4.65	892.25 891.47 890.64 889.19	6/29/83 7/18/83 11/30/83 12/13/84
BCC-5A	887.25	889.85	-6.10 -6.49 -6.92 -6.19 -5.39	881.15 880.76 880.33 881.06 881.86	6/29/83 7/19/83 9/1/83 11/30/83 12/12/84
ECC-6A	885.50	887.62	-4.45 -3.59 -3.12	881.05 881.91 882.50	9/2/83 11/30/83 12/12/84
ECC-7A	881.53	883.93	-8.50 <sup>b</sup> -2.43 -2.61	873.03 <sup>b</sup> 879.10 878.92	9/1/83 11/30/83 12/12/84
ECC-8A	885.42	886.22	-3.27	882.15	12/12/84
ECC-9A	881.01	883.11	+0.08	881.09	12/12/84
ECC-10A	879.60	882.30	-5.71	873.89	12/12/84
ECC-11A	884.40	886.90	-3.43	880.97	12/12/84

<sup>&</sup>lt;sup>a</sup>Positive sign indicates water level above ground surface; negative sign indicates water level below ground surface.  $^{\rm b}$  Noted while drilling with hollow stem augers.

GLT360/50-2

It is important to note that although data are scant, it appears that upward gradients in the shallow groundwater flow system occur beneath much of the site. In fact, the upper two hydrogeologic units may possibly act as separate aquifers in places. That is, the sand and gravel zone may be semiconfined in places due to lithologic variations in the upper saturated zone. Hydraulic gradients in the shallow flow system vary between approximately 0.01 ft/ft and 0.06 ft/ft. The actual gradients directly beneath the site are uncertain.

Water level data in the deep, confined aquifer indicate that flow is generally north to south. The maximum observed gradient in the deep confined aquifer was found to be 0.005 between wells ECC-3C and ECC-4C. Vertical gradients are upward since the potentiometric surface of the zone is above ground surface.

# Groundwater Contamination

Monitoring Well Results. The 15 monitoring wells at ECC were sampled in three phases during the RI. Samples were analyzed at the CLP for inorganics, volatiles, acids, base/neutrals, pesticides and PCB's.

Inorganic results from all three phases of sampling are presented in Table 4-11 for the shallow monitoring wells and

# TABLE 4-11 GROUNDWATER INDRSANIC RESULTS (ug/L) SHALLOW MONITORING WELLS ECC Site RI Report

		le Location: ite Sampled: ITR Number:	<b>97-19-83</b>	19-01 11-29-83 <b>HS09</b> 27	19-62 11-29-63 MS0928	1A-001 12-13-84 NE4629		29-801 87-19-83 MS8284	29-01 11-29-63 MS0930	29-001 12-13-64 NE4628	39-601 67-19-83 MS6265	3 <del>0-08</del> 2 07-19-83 MS0288	3A-01 11-29-63 MS0933	3A-881 12-13-84 NE4625	
COMPOUND	DETECTABLE LIMITS	QUALITY CRITERIA c		<del></del>											
ALLMINUM ANTIMONY ARSENIC	200 20 10	146 d			486	384				(65)	839	328		[12 <b>8</b> ]	
BARIUM BERYLLIUM	1 <b>00</b> 5	50 j 1000 j 0.0039 g	386	366	357	328		330	268	287	570	560	1078	868	
CADMIUM CALCIUM CHRONIUM CUBALT	1  10 50	10 j — 56 j	N/A	N/A	N/A	9577 <b>9</b> 11	E	N/A	N/A	98200 E 11	N/A 13	N/A	N/A	7 <b>8248</b> 15	
COPPER	59	1000 e												[163	
IRON LEAD CYANIDE	59 5 19	390 e 50 j 200 d		3870	3380	1454 6.7	••••	2748	3360	2931	8399	6330	18488	297	
NAGNESIUM NANGANESE	10	50 e	N/A	N/A 1 <b>6</b> 3	N/A 95	3466 <b>0</b> 66	Ε	N/A 56	N/A 49	32070 E	N/A 260	N/A 230	N/A 97	131 <b>999</b> 7 <b>9</b>	
MERCURY NICKEL POTASSIUM	9.2 49 —	0. 014 d 13. 4 d		b	• • • • • • • • • • •	• • • • • • • • • •	••••	<b>0.</b> 3 b	0.4	65	<b>0.</b> 3 42	b 77	89	84 1 <b>8</b> 5940	
SELENIUM SILVER	2 18	10 j 56 j		ස	14						3	4			
SODIUM THALLIUM TIN	 10 20	18 d	N/A	N/A	N/A	19969		N/A	N/A	15490	N/A	N/A	N/A	380700	
VANADIUM ZINC	200	5000 e	·	45	14	69			11	260			19	250	I

#### FOOTNOTES:

- a- QA data indicate the presence of these metal contaminants in the laboratory method blank
- b- This metal was also detected in the analysis of the field blank.
- c- U.S.EPA Drinking Nater Quality Criteria or National Drinking Nater Standards.
- d- Water Quality Criteria for Human Health Tocicity Protection (adjusted for consumption of mater only.)
- e- Secondary drinking water standard.
- g- Water Quality Criteria for Human Health U.S.EPA assigned carcinogen risk level of 10 -6 (adjusted for consumption of water only). One additional case of cancer in a population of 1,800,000
- h- No adverse effect level calculated by NAS/NRC.
- j- Primary drinking water standard.
- E- Value is estimated or not reported due to the presence of interference.
- R- Spike sample recovery is not within control limits.
- II- Positive values less than the contract required detection limit.
- N/A- Not analyzed for.
- Criteria has not been established for this compound.

# **TABLE 4-11** GROUNDMATER INDREGNIC RESULTS (ug/L) SHALLOW MONITORING WELLS EDC Site RI Report

	Da	e Location: te Sampled: ITR Number:	<b>87-19-83</b>	50-01 11-30-03 MS0936	5 <del>0-88</del> 1 12-12-84 NE4622	50-082 12-12-84 ME4639	6A-01 11-30-63 MS0937	6A-001 12-13-84 ME4627	7A-01 11-30-83 MS0938	7A-02 11-30-83 MS0939	79-001 12-13-84 ME4626	8A-001 12-13-04 HE4631	189-881 12-12-84 ME4624	Blank 07-19-83 MS8276	BLANK 11-38-83 MS8948	BLANK 12-13-84 ME4632
COMPOUND	DETECTABLE LINITS	GUALITY CRITERIA c														
ALIMINUM ANTIHONY ARSENIC	200 20 10	146 d 50 J	172 <b>0</b> 4	361		[140]		[66]	61500	663	[77]	[144]	[72]			(57)
BARIUM BERYLLIUM	190 5	9. 9639 g	398	392	413	438	598	612	875	397	331	353	296			
CADMILIN CALCIUM CHRONIUM CUBALT COPPER	1 10 50 50	18 ) 58 ) 1880 e	N/A 11	N/A	94898 13	99418 E 12	N/A	1611 <b>00</b> E	N/A 144 80 106	N/A	7355 <b>0</b> i	E 98586 E	779 <b>00</b> E	N/A	N/A	[900] E
IRON LEAD	59 5	300 e 50 j 200 d	7410	3290	5865	356	5478	1194	1 <b>05000</b> 102	1030	[73] 6. 5	2545	(51)	**********	210	[98]
CYANIDE Hagnesium Hanganese	1 <del>0</del> 10	50 e	N/A	N/A 52	33148 E 73	34160 E 50	N/A 231	6973 <b>8</b> E 94	N/A 1930	N/A 113	2978 <b>0</b> 57	E 38890 E 24	31448 E 40	N/A	N/A	[334] E
MERCURY NICKEL POTASSIUM	9.2 48 —	0.014 d 13.4 d			(32)			46 [2129]	<b>0.</b> 2 176		(2625)	(11 <b>95</b> )	[4765]	11.2	9.8	[34]
SELENIUM SILVER	2 1 <b>0</b>	10 J 50 J													20	
SCOTUM THALLIUM TIN	10 20	18 d	N/A 9. 4	N/A	10980	11219	N/A	118000	N/A	N/A	22388	151 <b>39</b>	25529	N/A 8. 4	N/A	1424
VANADILM ZINC	2 <b>00</b> 1 <b>0</b>	58 <b>00</b> e		36	155	158	35	42	276	31	37	69			49	31

### FOOTNOTES:

- a- QA data indicate the presence of these metal contaminants in the laboratory method blank b- This metal was also detected in the analysis of the field blank. c- U.S.EPA Drinking Mater Quality Criteria or National Drinking Mater Standards, d- Mater Quality Criteria for Human Health Tocicity Protection (adjusted for consumption of water only.)
- de nater visiting water standard.

  g- Nater Quality Criteria for Human health locality Protection (adjusted for consumption of water only). One additional case of cancer in a population of 1,800,800 he No adverse effect level calculated by NOS/NRC.

  j- Primary drinking water standard.

  E- Value is estimated or not reported due to the presence of interference.

  R- Spike sample recovery is not within control limits.

  (1) Positive values less than the contract required detection limit.

  NOS-Not analyzed for

- N/A- Not analyzed for.
- Criteria has not been established for this compound.

in Table 4-12 for the deep and intermediate wells. Two wells monitor the shallow saturated zone, well 3A and 11A. Background water quality is represented by wells 1A and 2A in shallow sand and gravel aquifer upgradient of the site. Inorganic analysis was not performed on well 11A samples due to very slow well recharge.

Inorganic constituents in well 3A found exceeding water quality standards or criteria and exceeding background levels in 1A and 2A are barium, iron, and nickel. Barium is only slightly above the primary drinking water standard of 1,000 ug/L. Iron is substantially above background though it is an aesthetic (taste) concern only. Nickel exceeded the EPA water quality criteria in well 3A although the background level in 2A also exceeded the criteria.

Inorganic constituents in the shallow confined aquifer found exceeding water quality standards or criteria and exceeding the background levels in wells 1A and 2A are:

- o aluminum in wells 5A and 7A
- o chromium in well 7A
- o iron in well 5A and 7A
- o lead in well 7A
- o nickel in 7A

**TABLE 4-12** GROUNDWATER INDREAMIC RESULTS (ug/L) DEEP & INTERMEDIATE MONITORING WELLS ECC Site RI Report

							DEE	P WELLS					INTERMEDI	ATE WELLS
		Samp	le Location: ate Sampled: ITR Number:	<b>87-18-63</b>	1C-01 11-29-63 MS0929	2C-901 07-18-83 MS0272	2C-01 11-29-63 MS0932	3C-061 07-18-83 MS0273	3C-01 11-30-84 MS0934	4C-881 87-18-83 MS8274	4C-002 07-18-63 MS0275	4C-01 11-30-83 MS0935	29-601 07-19-63 MS8271	29-01 11-29-83 MS0931
COMPOLIND	DE	TECTABLE LIMITS	QUALITY CRITERIA c											
ALUMINUM ANTIHUMY ARSENIC BARIUM BERYLLIUM		200 20 10 100 5	146 d 50 J 1600 J 8.0039 g	660	657	380	478	210	264	510	510	563	150	188
CADMIUM CALCIUM CHROMIUM CUBALT COPPER	N/A	1 18 50 50	10 J 50 J 1000 e		••••••	•		•••••	••••••	•	**********	•	•••••••	•••••
IRON LEAD CYANIDE HORNESTUM HONGONESE	N/A	50 5 10 	380 e 50 j 200 d 50 e	i	736 28	67 <b>0</b> 17	875 23	182 <b>0</b> 51	172 <b>0</b> 39	859	97 <b>9</b>	1 <b>989</b> 23	920 54	1140
MERCURY NICKEL POTASSIUM SELENIUM SILVER	******	0.2 46  2 10	9. 014 d 13. 4 d 18 J 58 J	• • • • • • • • • • • • • • • • • • •		••••••••••	@. 4 33	***************************************		42	25	19	<b>0.</b> 3 t	
SODIUM THALLIUM TIN VENADIUM ZINC	N/A	18 29 298 19	18 d		19	•••••••	26	••••	•••••••		••••••	74		••••••

#### FOOTNOTES:

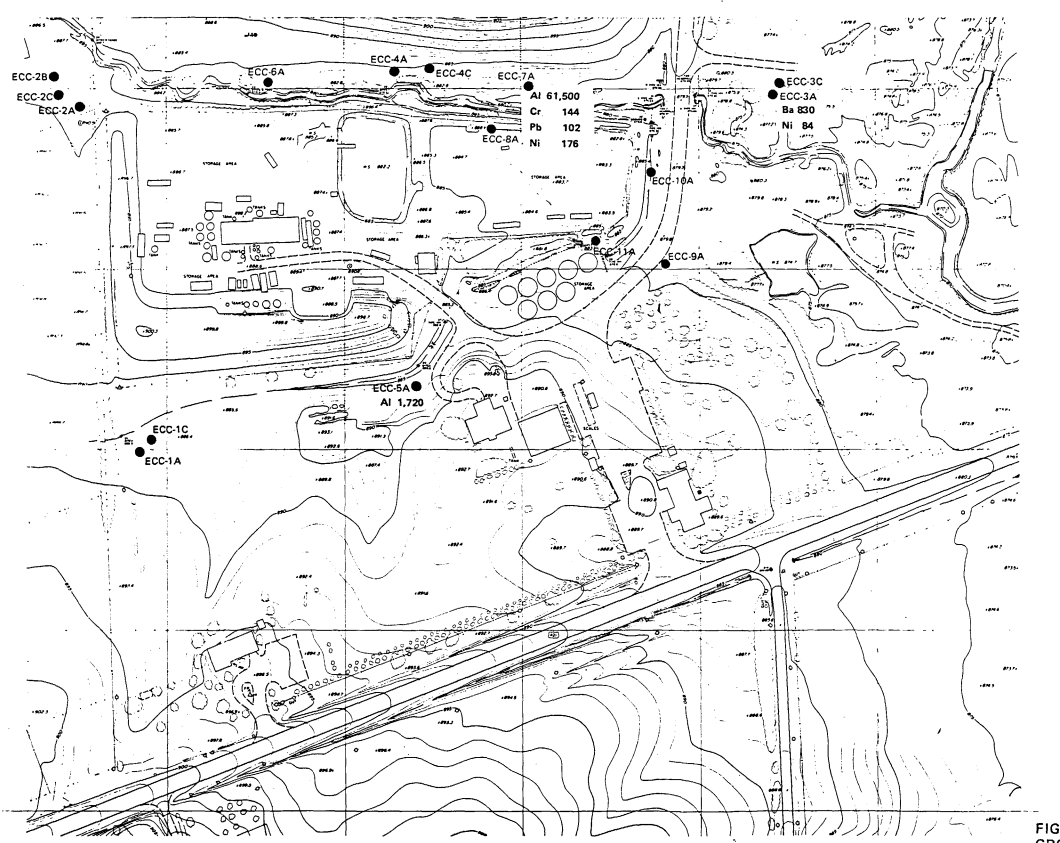
- a- 0A data indicate the presence of these metal contaminants in the laboratory method blank
  b- This metal was also detected in the analysis of the field blank.
  c- U.S.EPA Drinking Nater Quality Criteria or National Drinking Nater Standards.
  d- Nater Quality Criteria for Human Health Tocicity Protection (adjusted for consumption of water only.)
  e- Secondary drinking water standard.
  g- Nater Quality Criteria for Human Health U.S.EPA assigned carcinogen risk level of 10 -6 (adjusted for consumption of water only). One additional case of cancer in a population of 1,000,000.
  h- No adverse effect level calculated by NGS/NGC.
  1- Designed drinking water standard.
- 1- Primary drinking water standard. E- Value is estimated or not reported due to the presence of interference. R- Spike sample recovery is not within control limits. [1- Positive values less than the contract required detection limit.

- N/A- Not analyzed for.

   Criteria has not been established for this compound.

The aluminum levels in 5A (1,720 ug/L) and 7A (61,500 ug/L) exceed the EPA drinking water criteria of 73 ug/L. The background levels in 1A also exceed the criteria though not by the same extent. Barium is higher than background in well 7A, though it does not exceed the primary drinking water standard of 1,000 ug/L there. Chromium exceeds the primary drinking water standard of 50 ug/L in well 7A, where, it is 144 ug/L. The secondary water quality standard for iron was exceeded in all shallow wells, including background. In wells 5A and 7A levels were substantially higher than background. These levels are not a health threat. Lead was twice the primary drinking water criteria in well 7A where it was 102 ug/L. Nickel exceeded the EPA water quality criteria in well 7A as well as the background well 2A. Only in well 7A was it substantially higher than the background level. In summary, shallow wells 5A and 7A appear to have inorganic constituents in levels exceeding background that also exceed water quality criteria or standards. Figure 4-17 presents the distribution of inorganic constituents exceeding background levels and water quality criteria or primary drinking water standards.

In the deep confined aquifer inorganic constituents did not exceed background levels. Two inorganics, manganese and nickel, however, do exceed criteria or standards.



# LEGEND

REMEDIAL INVESTIGATION MONITORING WELL ECC-7A

NOTE: All well locations are approximate.

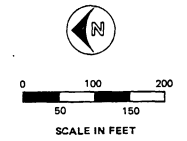


FIGURE 4-17
GROUNDWATER INORGANIC CONSTIUENTS EXCEEDING
BACKGROUND LEVELS AND WATER QUALITY
CRITERIA OR PRIMARY DRINKING WATER
STANDARDS (ug/l)
ECC RI

Organic results for the shallow monitoring wells are presented in Table 4-13 and for the deep and intermediate wells in Table 4-14. As discussed previously, wells 1A and 2A are representative of background water quality.

Several organics found in these wells and other shallow as well as deep wells are due to sampling bottle and/or laboratory contamination. Methylene chloride was found in nearly all samples and field blanks. It is used in preparatory cleaning of the VOA vials used for the samples. Acetone also was found in numerous samples as well as field blanks. Reagent grade acetone was used for equipment decontamination. Tetrachloroethene and trichloroethene were detected in wells 1A, 2A, and 5A at levels less than 9 ug/L quantification limit during the November 29-30, 1983 sampling. These levels are not considered to be representative of the groundwater since they were not detected in sampling phases before and after. Also wells 1A and 2A are upgradient of the site and would not be expected to show contamination.

Wells 3A and 11A monitoring the shallow saturated zone were found to be contaminated. Well 11A had high levels of trans-1,2-dichloroethene (4,000 ug/L) and trichloroethene (28,000 ug/L). Well 3A is contaminated with 13 VOC's. Compounds substantially above water quality criteria are benzene (<9 ug/L) 1,1-dichloroethane (96 ug/L), chloroform

#### TABLE 4-13 GROUNDWATER DRGANIC RESULTS (ug/L) SHALLOM MONITORING WELLS ECC Site RI Report

	Da	e Location: te Sampled: 0 ITR Number:	7-19-83	19 <del>-8</del> 1 11-29-83 \$28 <b>8</b> 3	19 <del>-62</del> 11-29-83 \$2801	1A-001 12-13-84 E7493	29- <b>00</b> 1 <b>07-</b> 19-83 \$2384	2 <del>9-0</del> 1 11-29-83 \$28 <b>8</b> 4	29-801 12-13-84 E7492	3A- <b>80</b> 1 87-19-83 523 <b>8</b> 5	3 <del>A-<b>88</b>2</del> 87-18-83 S2388	3 <del>9-8</del> 1 11-2 <del>9-</del> 83 528 <b>9</b> 7	3A- <b>86</b> 1 12-13-84 E7489	\$0-001 07-19-83 52386	5A-01 11-30-63 52810
VOLITILE COMPOUNDS	DETECTABLE LIMITS	QUALITY CRITERIA c	<del></del>	*****		~+ <del>**</del>		******		*****					
BENZENE 1, 1, 1-TRICHLORDETHONE 1, 1-DICHLORDETHONE CHLORDETHONE CHLORDETHONE CHLOROFORN	5 5 5 1 <b>0</b> 5	8.67 g 1986 d 8.94 g 								5 K 96 128	86 116	9 K 51.2 <del>18.</del> 7 9 K	4 J 1 <b>00</b>		
1, 1-DICHLOROETHENE TRANS-1, 2-DICHLOROETHENE TRANS-1, 3-DICHLOROPROPENE ETHYLEREZENE METHYLENE CHLORIDE	5 <b>5</b>	87 d 2466 d 6, 19 g	••••••	9 KE	9 КВ	22	************	11 B	3 J	19 8 B	16 8 K	9 K 77.5 9 K 18 B	1 <b>0</b> 3 J	••••••	9 KB
TRICHLOROF-LUOROMETHANE TETROCHLOROETHENE TOLUENE TRICHLOROETHENE VINYL CHLORIDE	5 5 5 5	8. 19 g 9. 8 g 15000 d 2. 8 g 2. 6 g	••••••	9 K	9 K	•••••••		9 K	***********	9 7	, , 6	9 K 9 K 85.8		•••••••	9 K 9 K
acetone 2-butanone Styrene Total Xylenes	i 5 i 5	990 h 1480 m		9649 B 9 K	9897 B		64 <b>8</b> 9	3016 B		1400	1400	1 <b>5030 B</b> 12	••••	490	54.5 B
TOTAL VOC's	J		0	27	18	8	9	18	8	256	231	320	117	8	18
BASE/NEUTRAL COMPOUNDS FLUORANTHENE ISOPHORONE	- 20 20	188 d 5569 d								218 K	218 K				
N-MITROSODI PROPLYAMINE BIS (2-ETHYLHEXYL) PHTHALATE DIETHYL PHTHALATE CHRYSENE PYRENE	20 20 29 20	21996 d 449969 d 9.9931 g 9.9931 g			23 K					29 K 29 K 30	: 218 K 318				
TOTAL BASE/NEUTRAL COMPOUNDS			8	•	23	0	0	8	0	110	116	9	9	8	8
TENTATIVELY IDENTIFIED COMPOUNDS	A														
1, 1-0XYBISETHANE 2-NETHYL-2-BUTANOL TETRAHYDROFURAN TRIPHENYLESTER PHOSPHORIC AC	- ID						4.2				13 4.2			5. 8 6. <b>0</b>	
***************************************		***********	****		E-F		125.553 <del>124</del> 865	**********					*********		

#### FOOTNOTES:

- A. Tentatively identified compound concentrations are estimated. A 1:1 response is assumed.

  B. Analyte has been found in the laboratory or field blank as well as the sample. Indicates probable contamination.

Applies to pesticide parameters where the identicication has been confirmed by 6C/MS.

- J. Indicates an estimated value. When mass spectral data indicates the presence of a compound that meets the identification criteria and the result is less than the specified detection limit but greater than zero.
- K. Actual value, within the limitations of the method is less than the value given
- N/A- Not analyzed for.
  - c- U.S. EPA Drinking Water Quality Criteria or National Drinking Water Standards.
  - d- Water Quality Criteria for Human Health Toxicity Protection (adjusted for consumption of mater only).

- g- Water Quality Criteria for Human health U.S.EPA assigned carcinogen risk level of 18 -6 (adjusted for comsumption of water only). One additional case of cancer in a population of 1,800,800. h- No adverse effect level calculated by NAS/NRC.
- i- Nonpriority hazardous substance.
- J- Total VOC's do not include the likely bottle and sampling contaminants methylene chloride and acetome, or other probable contaminants with footnote B.
- m- U.S.EPA 10-day health advisory level.

#### **TABLE 4-13** GROUNDWATER ORGANIC RESULTS (ug/L) SHALLOW MONITORING WELLS ECC Site RI Report

	ate Sampled:	12-12-84	59 <del>-08</del> 2 12-12-84 E7494	6 <del>9-8</del> 1 11-3 <b>8-</b> 83 52811	6 <del>A-00</del> 1 12-13-84 E7491	7A-81 11-38-83 52812	78-82 11-38-83 52813	7A <del>-88</del> 1 12-13-84 E7 <b>498</b>	8A-801 12-13-84 E7495	9 <del>0-00</del> 1 12-13-84 E7487	1 <b>8A-88</b> 1 12-12-84 F7488	11A-801 12-13-84 F7485	BLÂNK-001 07-19-83 \$2376	BLANK 11-39-83 52814	BLANK 12-13-84 E7496
DETECTABLE LIMITS	GUALITY														
5 5 5 19								4 J 98	7		29				
5					3 J	B • • • • • • • • • • • • • • • • • • •								*******	д 
5 5	87 d 2 <b>480</b> d		<b>4</b> J	19.5 R		16.5 R	9 кв	4.3	13	2 :	- '		34	9 R	
5 5 5 5	8, 19 g 8, 8 g 15 <b>888</b> d 2, 8 g				•	•						28000			• • • • • • • • • • • • • • • • • • • •
i 5	989 h		••••••	4284 B	24 B	23. 9 B 9 K	38.3 B		52 B	41 8	53 E 26 E		•••••	9 K	B 186
j		8		0	0	9	9	197	47	3	40	32900	0	21	16
20 28															
20 20 20 20	440000 d 8.0031 g	ı													99
*****************		0	8	0	9	0	9	9	0	9	0	9	0	0	95
A															
D													3. 8		
	DETECTABLE LIWITS  5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	Date Sampled: ITR Number:  DETECTABLE LIMITS CRITERIA c  5	S 8.67 9 1900 d 5 9.49 9 9 8.833 9 8 7 d 2480 d 5 8.19 9 3 J 5 8.8 9 5 15800 d 2.8 9 2 8 9 1 4 8 9 1	Date Sampled: 12-12-84	Date Sampled: 12-12-84 12-12-84 11-38-83 ITR Number: E7486 E7494 S2811  DETECTABLE GUALITY CRITERIA c  5 6.67 9 5 1900 d 5 0.94 9 10	Date Sampled: 12-12-84 12-12-84 11-38-83 12-13-84 ITR Number: E7486 E7494 S2811 E7491  DETECTABLE GUALTY LIMITS CRITERIA c  5	Date Sampled: 12-12-64 12-12-64 11-30-63 12-13-64 11-30-63 ITR Number: E7486 E7494 S2811 E7491 S2812  DETECTABLE QUALITY LIMITS CRITERIA c  5	Date Sampled: 12-12-84   12-12-94   11-38-83   12-13-94   11-38-83   11-38-	Date   Sampled:   12-12-04   12-12-04   11-30-03   12-13-04   12-13-04   12	Date Sampled: 12-12-64   12-12-64   11-36-83   12-13-64   11-36-83   12-13-64   12-13-64   11-36-83   12-13-64   12-13-64   11-36-83   12-13-64   12-13-	Date Sampled: 12-12-84   11-38-83   12-13-84   11-38-83   11-38-83   11-38-83   12-13-84   12-13-	Date Sampled: 12-12-94 11-39-83 12-13-94 11-39-83 12-13-94 12-13-1	Date Samples   12-12-04   12-12-04   12-13	Date Sampler: 12-12-64 12-12-64 11-38-63 12-13-64 11-38-63 12-13-64 12-13-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-64 12-13-13-64 12-13-64 12-13-64 1	Date Samples: 12-12-04 12-13-04 12-13-04 11-30-03 12-13-04 12-13-13-04 12-13-04 12-13-04 12-13-04 12-13-04 12-13-04 12-13-04 12-1

- A. Tentatively identified compound concentrations are estimated. A 1:1 response is assumed.
- B. Analyte has been found in the laboratory or field blank as well as the sample. Indicates probable contamination. C. Applies to pesticide parameters where the identicication has been confirmed by SC/MS.
- C. Applies to pesticide parameters where the identicication has been confirmed by GC/MS.

  J. Indicates an estimated value. When mass spectral data indicates the presence of a compound that meets the identification criteria
- and the result is less than the specified detection limit but greater than zero. K. Actual value, within the limitations of the method is less than the value given
- N/A- Not analyzed for.
- c- U.S. EPA Drinking Water Quality Criteria or National Drinking Water Standards.
- d- Water Quality Criteria for Human Health Toxicity Protection (adjusted for consumption of water only).
- e- Secondary Drinking Water Standard.
- g- Water Quality Criteria for Human Health U.S.EPA assigned carcinogen risk level of 10 -6 (adjusted for communition of water only). One additional case of cancer in a population of 1,000,000. h- No adverse effect level calculated by NAS/NRC.
- i- Nonpriority hazardous substance.
- j- Total VOC's do not include the likely bottle and sampling contaminants methylene chloride and acetone, or other probable contaminants with footnote B.
- m- U.S.EPA 19-day health advisory level.

# TABLE 4-14 GROUNDWATER ORGANIC RESULTS (ug/L) DEEP & INTERMEDIATE MONITORING WELLS ECC Site RI Report

INTERMEDIATE WELLS

DEEP WELLS

							DEC. III						
		ele Location: Pate Sampled: ITR Number:	<b>87-18-83</b>	10-01 11-29-83 \$28 <b>0</b> 2	2C- <b>86</b> 1 97-18-63 \$2372	20-01 11-29-83 \$28 <b>8</b> 6	3C- <b>89</b> 1 87-18-83 \$2373	3C- <b>6</b> 1 11-38-63 528 <b>6</b> 6	4C-891 97-18-83 \$2374	4C- <b>88</b> 2 97-18-83 52375	4C-01 11-30-83 \$2809	2 <b>B-801</b> <b>07-</b> 19-83 <b>S</b> 2371	28-01 11-25-83 \$2805
VOLATILE COMPOUNDS	DETECTABLE LIMITS	CRITERIA C	:						<del></del>				
richlordethane Ordethane Ne	5 5 5 18 5	0.67 s 1900 d 0.94 s	i										
.OROETHENE 2-DICHLOROETHENE 3-DICHLOROPROPENE ENE CHLORIDE	5 5	<b>9. 9.3</b> 3 9 87 0 24 <b>89</b> 6 <b>9.</b> 19 9	i i	9 +	ζ	9 #	 (B 5	K 12.4 1	3		9		9 1
OFLUGROMETHANE DROETHENE DETHENE LORIDE	5 5 5 5	2.8		••••••	•••••••	• • • • • • • • • • • • • • • • • • • •	••••••	•••••	••••••	•••••	••••••	•••••	•••••
ONE YLENES	5 5	998   1488	1	9 1	KB 1000 5			558, 7	··········		9	KB	9 }
)C¹s	)		• • • • • • • • • • • • • • • • • • •	0	5	9	8	8	0	0	0	0	8
EUTRAL COMPOUNDS  NE ILPROPLYAMINE LHEXYL) PHTHALATE ITHALATE	29 28 29 29 26 28	21999 449999 9.9931	d d g										
rse/neutral compounds			(	0	0	0	0	9	0	9	8	9	
TENTATIVELY ENTIFIED COMPOUNDS BISETHANE L-2-BUTANDL DROFURAN VLESTER PHOSPHORIC ACID	4												
8. C. J. K N/A- c- e- 9-	Analyte has Applies to p Indicates an And the resu Actual value Not analyzed U.S.EPA Drin Water Qualit Moadverse e	been found i esticide par estimated v ult is less t is within the for. kking Water G inking Water y Criteria f	n the labo ameters what alue. When han the so limitation uality Crior Human Standard or Human calculate	oratory or fi here the iden n mass spectr pecified dete ons of the me iteria or Nat Health - Toxi	eld blank as ticication half data indi- ection limit thod is less tional Brinki city Protect	ed. A 1:1 resided as the mell as the process the property of the process than the valing Water Station (adjusted carcinogen	sample. Indirmed by GC, esence of a than zero. lue given ndards. d for consu	ficates proba MS. compound tha aption of wat	t meets the	identificati		). Ome additi	ional case of

3- Total VOC's do not include the likely bottle and sampling contaminants methylene chloride and acetone, or other probable contaminants with footnote 8. m- U.S.EPA 18-day health advisory level.

(<9 ug/L), 1,1-dichloroethene (10 ug/L), trans 1,
3-dichloropropene (77.5 ug/L), trichloroethene (9 ug/L), and
vinyl chloride (85.8 ug/L). Well 3A also contained five
base/neutral compounds, one of which, pyrene, was
quantifiable. Pyrene was found at 30 ug/L, substantially
higher than the water quality criteria.</pre>

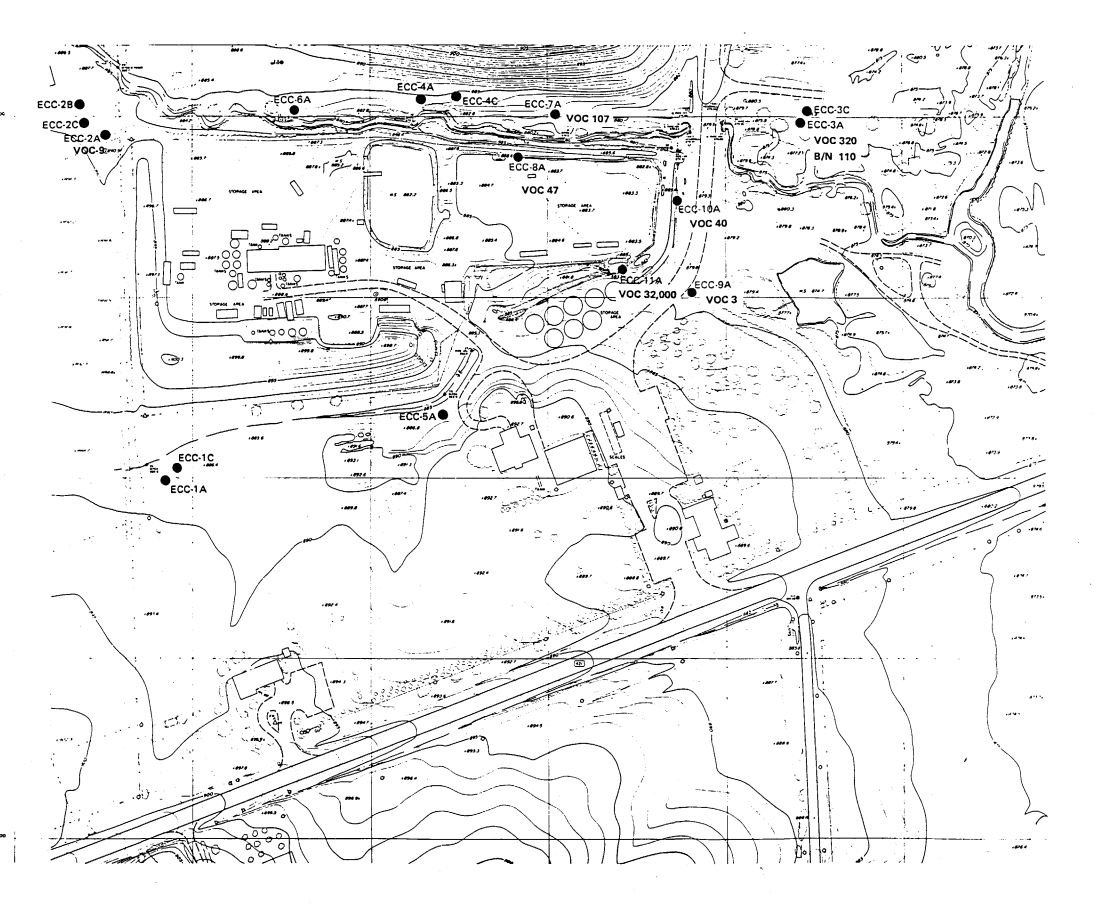
Organic groundwater contaminants in the shallow sand and gravel aquifer were found in wells 7A, 8A, 9A, and 10A. The following VOC's were most significant:

- o 1,1-dichloroethane, well 8A
- o chloroethane, wells 7A and 10A
- o 1,1, dichloroethene, wells 7A, 8A, and 10A
- o trichloroethene, well 8A

Figure 4-18 presents the distribution of total volatile organics and total base/neutrals.

No organic groundwater contaminants were found in the wells monitoring the deep confined aquifer.

In summary, the greatest organic contamination was found in the shallow saturated zone at well 11A, with lesser amounts at well 3A. The shallow sand and gravel aquifer was found to be contaminated at wells 7A, 8A, and 10A. The deep aquifer was not found to be contaminated.



#### LEGEND

REMEDIAL INVESTIGATION MONITORING WELL ECC.7A

VOC - TOTAL VOLATILE ORGANICS (ug/I)
B/N - TOTAL BASE/NEUTRALS (ug/I)
(NOT INCLUDING BLANK, FIELD OR
LABORATORY INDUCED CONTAMINATION)

NOTE: All well locations are approximate

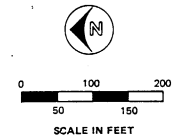


FIGURE 4-18
GROUNDWATER TOTAL VOLATILE
ORGANICS AND TOTAL BASE/NEUTRAL
CONCENTRATIONS (ug/l)
ECC RI REPORT

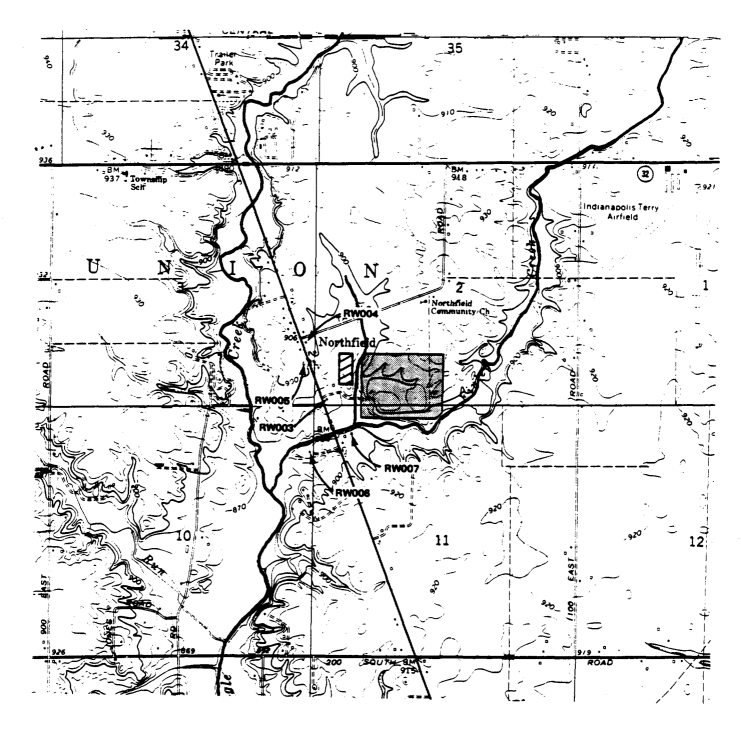
Residential Well Results. Six residential wells were sampled and analyzed for the full CLP inorganics and organics data packages (Figure 4-19). Inorganic results are presented in Table 4-15. Quality assurance review of laboratory data found reliability of the inorganic analysis to be strongly suspect and not considered useable. As discussed in Chapter 3, however, previous analysis of residential well samples has not found inorganics exceeding water quality standards with the exception of one sample at the Jennings well with lead at 93 ug/L.

Organic contamination was not found in any residential wells although acetone was reported in one sample, likely introduced during sampling.

### CONCLUSIONS AND OBSERVATIONS

Onsite soil investigations showed soil to be heavily contaminated, primarily with organic contaminants. Results of the hydrogeologic investigation have shown the existence of four hydrogeologic units in the area, a shallow saturated zone, a shallow sand and gravel aquifer, a clayey silt and silty clay zone, and a deep confined aquifer.

Migration of soil contaminants to the shallow saturated zone has occurred onsite as evidenced by high levels of contaminants in well 11A. Further leaching of soil contaminants to



#### LEGEND



NORTHSIDE SANITARY LANDFILL

ECC SITE



**FIGURE 4-19** RESIDENTIAL WELL SAMPLING LOCATION ECC RI REPORT

TABLE 4-15
RESIDENTIAL WELL INORGANIC RESULTS (ug/L)
ECC Site RI Report

	Samp	le Location: Well Name:	R <b>W88</b> 3 Bankert	RM994 Rousch	R <b>U00</b> 5 Jennings	RM <b>00</b> 5(duplicate) Jennings	RM006 Holly	RW007 Vandergriff	BLANK
COMPOUND	DETECTABLE LINITS	CRITERIA C							
ALIMINUM ARTIMONY ARSENIC BARIUM BERYLLIUM	280 29 19 100 5	146 d 50 J 1000 J 0.0039 g	482i 25 (9)	447i 28 [5, 5]	[66] i (7) 303	131i • 23 •	[97] i [7] 278	498i 24 [2,4]	486i 18 4
CADMILM CALCIUM CHROMIUM COBALT COPPER	1 19 59 59	18 J 50 J 1888 e	325i *	41 <b>6</b> i + +	1 <b>03900</b> [3.6] [42]	348i #	572 <b>66</b> [8, 9]	171i E10.33	44 (4.5) (3)
IRON LEAD CYANIDE MAGNESIUM MAGNESE	50 5 10 10	388 e - 58 J 288 d  58 e	[14] # 220	[9.2] # # 480	3290 6.0 # 40906 133	[11] * * 245	111 <b>0</b> ± 262 <b>00</b> 33.9	290 *	(39) * * *
NERCURY NICKEL POTASSIUM SELENIUM SILVER	0.2 40 - 2 18	8.014 d 13.4 d 	# [7] N/A #	[11] N/A +	# [16] N/A #	(7. 83 N/A	(19.3) N/A (7.7)	[8] N/A # #	† † N/A †
SODIUM THALLIUM TIN VANADIUM ZINC	10 20 200 18	18 d 	381 <b>800</b> i + + +	380000 # # #	153 <b>00</b> # # # 134	3638 <b>00</b> # # #	313 <b>90</b> 1 # # 49.2	25 <b>0000</b> # # #	143000 # # # #

#### FOOTNOTES:

- a- QA data indicate the presence of these metal contaminants in the laboratory method blank
  b- This metal was also detected in the analysis of the field blank.
  c- U.S.EPA Drinking Mater Quality Criteria or National Drinking Mater Standards.
  d- Mater Quality Criteria for Human Health Tocicity Protection (adjusted for consumption of mater only.)
  e- Secondary drinking water standard.
  g- Water Quality Criteria for Human Health U.S.EPA assigned carcinogen risk level of 18 -6 (for consumption of water only). One additional case of cancer in a population of 1,880,800.
  h- No adverse effect level calculated by NAS/NRC.
  i- Value has been corrected for the amount of contaminant in the lab blank.
  i- Deligant delighting mater standard.

- 1 Primary drinking water standard.
  E- Value is estimated or not reported due to the presence of interference.
  R- Spike sample recovery is not within control limits.
  []- Positive values less than the contract required detection limit.

- N/A- Not analyzed for.
- \* Less than laboratory detection limit (laboratory did not specify the limit)
- Criteria has not been established for this compound.

the saturated zone is expected to be greatly slowed due to the presence of a compacted clay cap on the northern half of the site and the continued existence of the concrete pad on the south half of the site.

The shallow sand and gravel aquifer has been shown to be contaminated with inorganics and organics in well 7A and lesser amounts of organics in wells 8A and 10A. Because of the presence of the NSL site east of ECC, it cannot be definitively stated that the source of contamination in wells 3A and 7A is ECC though the contaminants are consistent with those found onsite. Organic contamination in wells 8A and 10A is likely due to onsite soils at ECC since they are directly downgradient of ECC contaminated soils and not NSL.

Contamination of the shallow sand and gravel aquifer may have occurred either via migration through the silty clay till onsite or through contaminated water and sediment in the former cooling water pond. As discussed perviously, the cooling pond intersected the shallow sand and gravel aquifer.

The deep confined aquifer below the site has not been found to be contaminated. Future migration of onsite contaminants to the deep aquifer is highly unlikely due to the upward vertical hydraulic gradient. Migration of contaminants to the nearest residential wells surrounding the site is not indicated by the results of the residential well sampling.

# SURFACE WATER AND SEDIMENTS

A well-developed drainage pattern exists in the area surrounding the ECC site. The principal surface drainage features are Eagle Creek and Finley Creek, an associated tributary. Two minor surface drainage features are adjacent to the site. An unnamed ditch flows south along the eastern site boundary and converges about 1,000 feet downstream from the site with Finley Creek. The other unnamed ditch flows southeast along the western and southern site boundaries before discharging near the southeast corner of the site, into the unnamed tributary of Finley Creek. Finley Creek converges with Eagle Creek about one-half mile southwest of the site. Eagle Creek then flows south for about 10 miles before discharging into the Eagle Creek Reservoir. The site is located outside the 100-year flood plain. Enclosed in Appendix C of this report are aerial photographs and a topographic map illustrating the area surrounding the ECC and NSL sites.

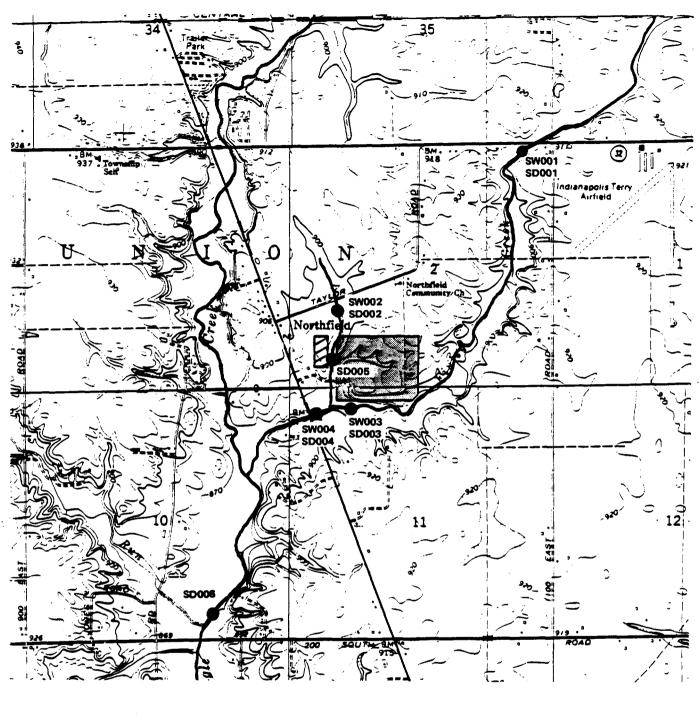
Natural surface water runoff from the area surrounding the site flows toward the unnamed tributary of Finley Creek or toward Finley Creek. The ECC site has been capped with clay

as part of the surface cleanup activities. Surface water runoff from the northern part of the site largely flows south where a berm along the edge of the concrete pad redirects runoff west to the ditch. Runoff from the concrete pad flows south and is routed through a pipe at the southeast corner of the site and to the unnamed ditch. Before capping, runoff was directed to the cooling pond and occasionally overflowed to the unnamed ditch.

## SCOPE AND METHODS

The purpose of the initial surface water and sediment sampling effort was to determine the extent of contamination in the unnamed ditch (east of the site), Finley Creek, and Eagle Creek. Previous ISBH and USGS sampling efforts have demonstrated contamination of surface water and sediment downstream from the ECC and NSL sites as shown in Chapter 3.

Four surface water samples and six sediment samples were taken on July 18, 1983, at locations in the unnamed ditch and Finley Creek identified in Figure 4-20. Surface water samples were collected at mid-depth of the stream with stainless steel dippers. Sediment samples were a composite of 6 to 14 cores from 1 to 3 inches long taken within a 10 foot square area. Details of sampling methods are described in Appendix A.



LEGEND

NORTHSIDE SANITARY LANDFILL

ECC SITE

SURFACE WATER AND SEDIMENT SAMPLING LOCATIONS (APPROXIMATE)



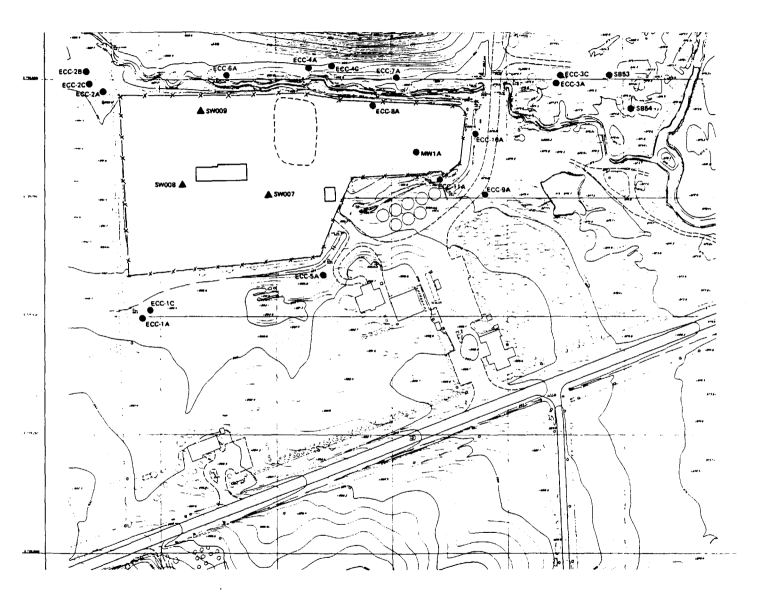
**FIGURE 4-20 SURFACE WATER AND SEDIMENT SAMPLING LOCATIONS** ECC RI REPORT

Three onsite surface water samples were collected on December 12, 1984, during the Phase 3 monitoring well sampling when sampling team members observed visibly contaminated water ponding on the clay cap onsite. The samples were collected from small areas of ponded water in the north half of the site (Figure 4-21). The site had been capped with 1 foot of clay previously. Sample bottles were filled by immersing in the ponded water. Inorganic samples were field filtered prior to preservation.

# RESULTS

Surface water and sediment samples were analyzed for concentrations of pollutants in four categories: inorganics, volatile organics, base/neutrals, and pesticides and PCB's.

A summary of inorganic results for the surface water samples is presented in Table 4-16 and a summary of inorganic results for sediments is presented in Table 4-17. Inorganic surface water data show elevated concentrations of aluminum, iron and manganese at SW-002 in the unnamed ditch upstream of the ECC and NSL sites. All three of these constituents are at levels above water quality criteria or standards at this location as well as downstream of ECC and NSL at SW-003 and SW-004. Manganese was also found at elevated levels at all 3 onsite sample locations. Mercury was found at SW-003 and SW-004 though detection in the field blank indicates it



LEGENO

ECC-7A REMEDIAL INVESTIGATION MONITORING WELL

▲ SURFACE WATER SAMPLE LOCATION

NOTE: All well locations are approximate

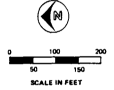


FIGURE 4-21
ONSITE SURFACE WATER
SAMPLE LOCATIONS
ECC RI REPORT

Table 4-16
INORGANIC ANALYTICAL RESULTS
SURFACE WATER SAMPLES
ECC SITE (ug/L)

( ′

		<u>.</u> .							
	SW-001	SW-002	SW-003	SW-004-01	SW-004-02	Blank	SW-007	SW-008	SW-009
	7/18/83	7/18/83	7/18/83	7/18/83	7/18/83	7/18/83	12/12/89	12/12/89	12/12/89
Compound									
Aluminum		3,050	340	<b>4</b> 90	440		[69]	[71]	[80]
Chromium		·							15
Barium					180		[108]	[92]	[173]
Beryllium									
Cobalt									
Copper									[13]
Iron	280	4,460	890	1,410	1,420		[80]	[77]	[83]
Nickel	47	·					[21]		
Manganese b	190	580	76	130	130		1,708	1,534	1,300
Zinc							36	41	79
Zincb Boron									
Vanadium								[9.2]	
Silver									
Arsenic									
Antimony									
Selenium			6						
Thallium <sup>a</sup>			_	_	_				
Mercury			0.2 <sup>C</sup>	0.4 <sup>C</sup>	0.3 <sup>C</sup>	0.2			
Mercury Tin									
Cadmium									
Lead									
Cyanide	0.007	0.013	0.005	0.008	0.013				

a QA data indicate poor or marginal recovery of these spiked metals.

QA data indicate the presence of these metal contaminants in the laboratory method blank.

 $<sup>^{\</sup>mathbf{C}}$ This metal also detected in the analysis of the field blank.

<sup>[] -</sup> positive values less then the contract required detection limit. blank = not detected.

Table 4-17
INORGANIC ANALYTICAL RESULTS
SEDIMENT SAMPLES
ECC SITE

	į	We'			SD-004			
Compound	SD-001	SD-002	SD-003	SD-004	(Duplicate)	SD-005	SD-006	Blank
Aluminum	2,172	9,744	4,326	2,890	3,050	5,928	2,850	200
Chromium	4	13	13	5	6	11	<5.7	9
Barium	45	102	44	31	35	66	27	<5
Beryllium	<0.45	0.6	<0.48	<0.25	<0.25	<0.57	<0.57	<0.25
Cobalt	<4.5	<12	5.3	<4	16	<9	<5,7	<2.5
Copper	7	23	19	13	16	23	9	<2.5
Iron	8,598	18,624	12,415	8,900	8,080	18,696	9,257	120
Nickel	<4	21	13	11	10	23	<11	<2
Manganese	161	499	275	170	158	413	239	<0.75
Zinc	<29	75	52	33	39	64	<30	<2
Boron	<9	<10	<10	<5	<b>&lt;</b> 5	<11	<11	<b>&lt;</b> 5
Vanadium	<18	23	<19	<10	<10	<23	<23	<18
Silver	<0.9	<1	<1	<0.5	<0.5	<1.1	<1.1	<0.5
Arsenic	<0.9	<1	<1	<0.5	<0.5	<1.1	<1.1	<0.5
Antimony	<2	<2	<2	<1	<1	<2	<2	<1
Selenium	<0.2	<0.2	<0.2	<0.1	<0.1	<0.2	<0.2	<0.1
Thallium	<0.9	<1	<1	<0.5	<0.5	<1.1	<1.1	<0.5
Mercury	<0.02	<0.02	2.25	<0.01	0.02	0.05	0.05	0.04
Tin	<2_	<2	<2_	<1_	<1_	<2	<2_	<1_
Cadmium	1.65°	2.3	1.83 <sup>C</sup>	0.82 <sup>C</sup>	0.78 <sup>C</sup>	1.41°	1.30 <sup>C</sup>	0.26°
Lead	19.0	11.5	31.3	17.5	32.3	48	6.8	1.9 <sup>c</sup>
Cyanide	33	<19	38	<10	196	73	<23	<10
Percent Moisture	45	48	48	_b	_p	56	44	_b

<sup>&</sup>lt;sup>a</sup>Concentrations expressed as mg/kg per dry unit weight except SD-004 and SD-004 duplicate.

b
Sample quantities were insufficient to determine moisture content.

<sup>&</sup>lt;sup>C</sup>QA review indicates that these data should be regarded as qualitative indication of the presence of these metals because the concentrations are below the lowest quantitative standard.

to be a sampling or laboratory contaminant. In summary, inorganic results do not show contamination of offsite surface water from either ECC or NSL at the locations sampled. Onsite inorganic contamination is limited to manganese.

Sediment inorganic results downstream of ECC showed only lead at concentrations above upstream levels. Lead was 48 mg/kg at SD-005 in the unnamed ditch whereas upstream of ECC and NSL it was 11.5 mg/kg. At location SD-005 the contaminant source could either be ECC or NSL.

Organic results are summarized in Tables 4-18 and 4-19 for the surface water and sediments, respectfully. Organic contamination of offsite surface water was limited to location SW-004. Compounds found at concentrations exceeding quantification limits were chloroethane, 1,1-dichloroethane, 1,1-trichloroethane, vinyl chloride, trans-1,2-dichloroethane and trichloroethene. Only 1,1-dichloroethane, vinyl chloride, and trichloroethene exceeded EPA water quality criteria at the 10<sup>-6</sup> cancer risk level.

Five tentatively identified organic compounds were also found in SW-004, though only one compound was confirmed in the duplicate sample.

Table 4-18
ORGANIC ANALYTICAL RESULTS (UG/L)
SURFACE WATER SAMPLES
ECC SITE

Compound	SW-001 7/18/83	SW-002 7/18/83	SW-003 7/18/83	SW-004-01 7/18/83	SW-004-02 7/18/83	Blank 7/18/83	SW-007 12/12/89	SW-008 12/12/89	SW-009 12/12/89
Base/Neutral Compounds <sup>a</sup>							_		
4-chloro-3-methyl phenol phenol 2-methyl phenol 4-methyl phenol bis(2-ethylhexyl)phthalate di-n-octyl phthalate isophorone		< 20					30 <sup>C</sup> 92 27 89	87	120 17 <sup>C</sup> 86
<u>Volatiles</u> <sup>b</sup>									
1,1,1-trichloroethane 1,1-dichloroethane chloroethane 1,2-transdichloroethene methylene chloride tetrachlorethene trichloroethene vinyl chloride ethyl benzene toluene acetone 2-butanone total xylenes	< 5	< 5	< 5	120 45 12 330 < 5 < 5 67 10	110 45 12 330 < 5 < 5 68 11	3,100	34 83 29 240 13 <sup>C</sup> 82 1,100 560 47	6 <sup>C</sup> 86 18 160  26 220 150 16 <sup>C</sup>	6 3 <sup>C</sup> 5 <sup>C</sup> 13 2 <sup>C</sup> 6 30 16
Nonpriority Pollutants/ Hazardous Substances									
o-xylene				< 5	< 5				
Tentatively Identified Compounds									
<pre>1,1,1-trichloro-1,2,2-trifluc trichloroethene 2,3,4-trimethylhexane 2,4-dimethylheptane 1,4-dioxane</pre>	oroethane			6.9 14 22 10	13	14			
tetrahydrofuran						7.1			

aQA review identified base/neutral results of 7/18/83 samples as semiquantitative because the average surrogate recovery is <40 percent. bQA review identified the volatile results of 7/18/83 samples acceptable due to good QA analytical results despite the fact that the analyses cwere run after expiration of the acceptable holding period. Indicates an estimated value.

Blank = not detected.

Table 4-19 ORGANIC ANALYTICAL RESULTS SEDIMENT SAMPLING ECC SITE

Compound <sup>a</sup>	SD-001	SD-002	SD-003	SD-004 <sup>b</sup>	SD-004 <sup>b</sup>	CD OOF	CD 000	<b>7</b> 1.5 - 1.5
Сомроина	<u>20-001</u>	SD=002	SD-003	<u>SD-004</u>	(Duplicate)	SD-005	SD-006	Blank
Base/Neutral Compounds								
bis(2-ethylhexyl)phthalate				C		912		
benzo(a)anthracene				440 <sup>C</sup>				
benzo(a)pyrene				< 800°				
benzo(b)fluoranthene				< 800 <sup>°</sup>				
benzo(k)fluoranthene				< 800 <sup>C</sup>				
chrysene				440 <sup>C</sup>				
benzo(ghi)perylene				< 800°C				
dibenzo(a,h)anthracene				< 800 <sup>C</sup>				
indeno(1,2,3-cd)pyrene				< 800°				
Volatiles								
methylene chloride	< 4.5	< 4.8	6.1	2.5	< 3	9.1	< 4.4	< 3.6
fluorotrichloromethane		< 4.8						
Nonpriority Pollutants/								
Hazardous Substances								
benzoic acid				< 4,000				
4-methylphenol				960	680			
TENTATIVELY IDENTIFIED COMPOUNDS								
Base/Neutral Compounds								
dichloromethane							170	
2-methyl-1-pentene							860	
1,3-dimethylbenzene			310				000	
•								

<sup>&</sup>lt;sup>a</sup>Concentrations expressed as ug/kg per dry unit weight except SD-004 and SD-004 duplicate.

Concentrations reported per wet unit because sample quantities were

insufficient to determine dry unit weight.

C Base/neutral analysis results were determined to be semiquantitative due to low recoveries in surrogate samples.

Three additional compounds (methylene chloride, o-xylene, and tetrachloroethene) were detected in surface water samples; however, concentrations were below quantifiable limits. Contamination of samples by methylene chloride is probably due to sample bottle contamination. Bis (2-ethylhexyl) phthalate was also detected in the upstream sample SW-002, but only in concentrations below the quantifiable limit.

Onsite ponded water was found to be contaminated with volatile and base/neutral compounds. All 3 locations showed contamination with location SW007 showing higher levels and more compounds. Several of the volatile compounds had also been detected at the offsite location SW-004. These were 1,1,1-trichloroethane, 1,2-transdichloroethene, tetrachloroethene and trichloroethene.

ECC site records and chemical analysis data implicate the ECC site as the source of contaminants identified at location SW-004. ECC site records report that chlorinated hydrocarbon solvents were processed at the facility. Further, drainage patterns direct over land flow from the vicinity of the ECC and NSL site towards sampling location SW-004. Sampling location SW-003 is approximately 750 feet upstream of location SW-004 on Finley Creek but receives runoff only from the NSL site. Surface water from this

sampling location was not found to be contaminated by chlorinated hydrocarbons.

Organic sediment contaminants were limited primarily to the base/neutral and acid fractions. Contaminants above the quantifiable limit are:

- o methylene chloride at all locations
- o bis (2-ethylhexyl) phthalate at SD-005
- o benzo(a) anthracene at SD-004
- o chrysene at SD-004
- o 4-methyl phenol at SD-004

Methylene chloride appeared in all samples including the blank and may be a sample bottle contaminant. SD-004 contaminants were not found in the duplicate sample with the exception of a 4-methyl-phenol. The base/neutral contaminants found at SD-004 were not found in any of the Phase 1 or 2 onsite soil samples. As a result it is not believed that ECC is the source of this potential contamination.

# CONCLUSIONS AND OBSERVATIONS

From the analysis of these results, the following conclusions are drawn:

- o Surface water runoff from the ECC site is directed towards the unnamed tributary of Finley Creek or towards Finley Creek.
- o Inorganic contamination of surface water does not appear to be occurring offsite in the vicinity of ECC.
- o Inorganic sediment contamination in the vicinity of ECC is limited to lead in the unnamed ditch.
- o Organic contamination of offsite surface water is limited to location SW-004. Contaminants consist almost entirely of chlorinated hydrocarbons.
- o Surface water ponded on the clay onsite was found to be contaminated with a variety of base/neutral and volatile compounds.
- o ECC site records and chemical analysis data implicate the ECC site as a source of organic contaminants detected in location SW-004.
- o Organic contamination of sediments possibly resulting from the ECC site was found at SD-005 (bis(2-ethylhexyl)phthalate) in the unnamed ditch and SD-004 in Finley Creek (4-methylphenol).

# Chapter 5

#### CONTAMINANT MIGRATION AND FATE

#### INTRODUCTION

This chapter is divided into two sections: general discussion of the present and potential pathways of contaminant migration in terms of the possible receptors; and a discussion of the migration and fate of contaminants at the ECC site. Due to the large number of contaminants found onsite, specific indicator chemicals were chosen as representative of the range of contaminants based on concentration, migration potential, degradation rates, toxicity, and carcinogenicity. The indicator chemicals chosen are listed in Table 5-1.

#### POTENTIAL PATHWAYS OF MIGRATION

## CONTAMINANT SOURCE

As a result of initial remedial measures, the original sources of contamination at the ECC site have been eliminated. The current source at the site is the subsurface soil which contains high concentrations of organic compounds as described in Chapter 4.

# Table 5-1 INDICATOR CHEMICALS AT ECC

Chloroform

Methylene Chloride

1,1,2-Trichloroethane (1,1,2 TCA)

1,1,1-Trichloroethane (1,1,1 TCA)

Trichloroethene (TCE)

Tetrachloroethene (PCE)

Ethylbenzene

Toluene

Phenol

PCB's

Bis(2-ethylhexyl)phthalate

Di-n-butyl phthalate

Diethyl phthalate

Dimethyl phthalate

GLT301/58

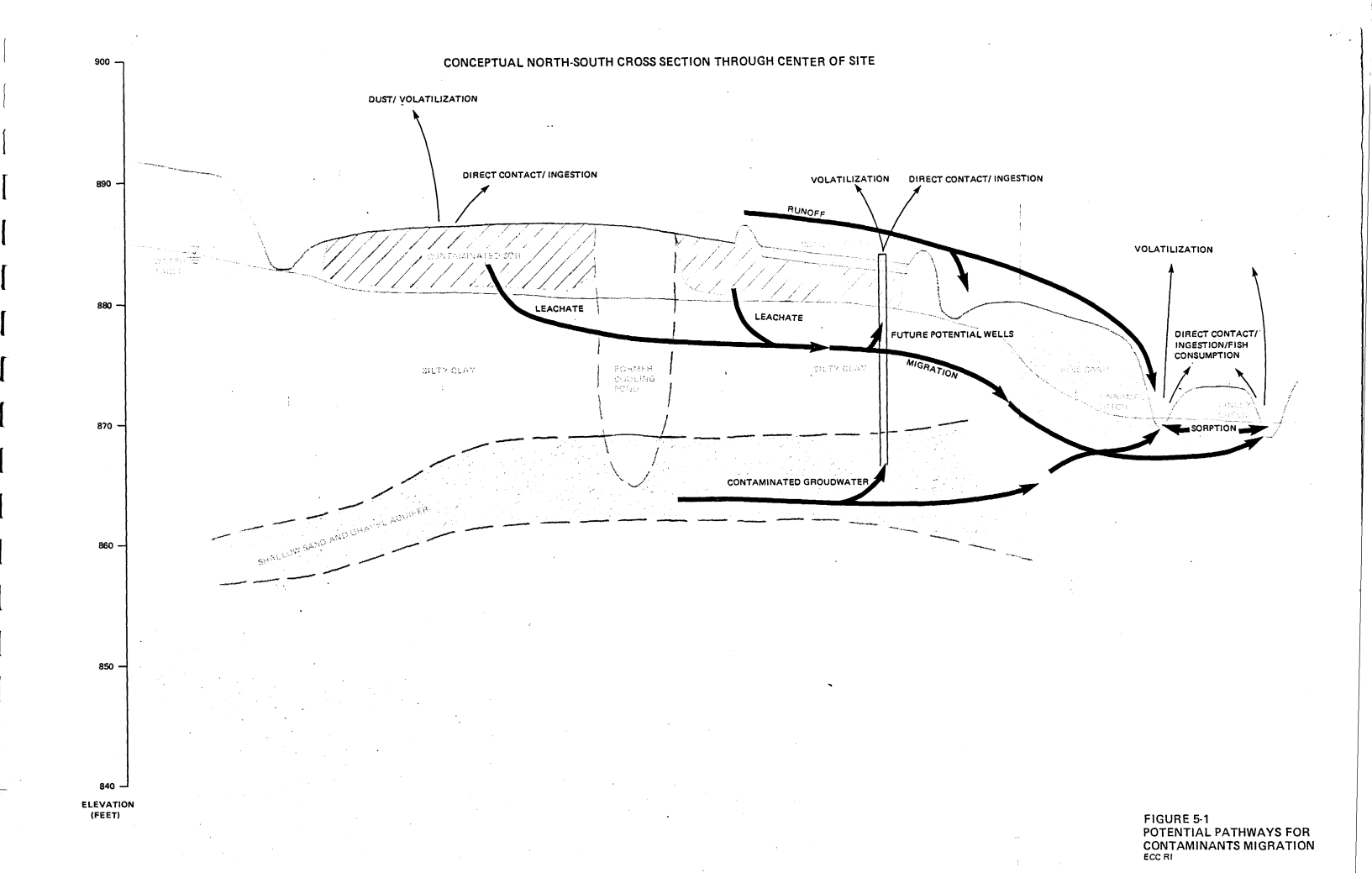
#### **PATHWAYS**

Figure 5-1 illustrates the potential pathways for contaminant migration.

#### Onsite Soils

Although the ECC site was covered with a clay cap upon completion of surface cleanup activities, samples from ponding surface water indicated the presence of organics. Though soil samples of the cap were not analyzed, it is presumed they are contaminated with the organics detected in the ponding surface water samples. These contaminants could volatilize or be transported as dust particles entrained by wind or transported in surface water runoff. Below the cap, heavily contaminated soil could be a risk to receptor populations since any future excavation might bring contaminants to the surface. Once chemicals are at the surface, receptors (plants and wildlife, as well as humans) may be subject to inhalation, ingestion and direct contact of harmful compounds.

Transport of contaminants from onsite soils is also likely through leaching. As water infiltrates through the contaminated soil, it will desorb many compounds and eventually leach into the groundwater in the shallow saturated zone. This is presently the case as the



groundwater samples from the shallow saturated zone were found to be contaminated with volatile organics.

# Groundwater

Once contaminants have entered the groundwater, several pathways of migration are possible. As mentioned previously in this report, four hydrologic units are located under the ECC site. In the past, contaminants could potentially migrate downwards from the shallow saturated zone and contaminate the lower sand and gravel aquifer. Low level contamination found in the shallow sand and gravel aquifer onsite indicates that this probably has occurred. Alteration of site characteristics during surface cleanup, however, has made this an unlikely migration pathway presently or in the future. The cooling pond, which was hydrologically connected to the shallow sand and gravel aquifer, has been cleaned of contaminated water and sediments, and backfilled with clean fill material. Onsite ponding water has also been removed, thus eliminating the downward vertical gradient. Water can no longer pond onsite, and vertical gradients between the shallow saturated zone and the shallow sand and gravel aquifer are upward. However, future excavation at the site could cause ponding of water onsite and reverse the gradient and enable downward migration of contaminants into the shallow sand and gravel aquifer.

Evidence of downward migration of contaminants from the shallow sand and gravel to the deep confined aquifer was not found and is highly unlikely now or in the future due to the upward vertical gradient. Existing low level contaminants in the shallow sand and gravel aquifer will likely migrate south and discharge to the unnamed ditch or Finley Creek. Receptors could potentially contact the groundwater if potable wells are constructed within the zones of contaminantion.

In summary, the most probable pathways for contaminant transport in the groundwater are through migration from the shallow saturated zone or from the shallow sand and gravel aquifer to the unnamed ditch or Finely Creek.

# Surface Waters

Both the unnamed ditch and Finley Creek receive groundwater and surface water runoff from the ECC site. Contaminants in the surface water may volatilize, precipitate or adsorb in sediments, or remain in solution and be transported downstream to Big Eagle Creek and eventually the Eagle Creek Reservoir. Receptors may be exposed by wading in the creek, ingesting contaminated water, or ingesting fish which have bioaccumulated contaminants.

## Sediments

Contaminants within stream sediment may dissociate and reenter solution or may be scoured and resuspended in high flow and carried downstream. During low flow periods contaminated sediments may be exposed along the stream banks and may be transported as dust.

# MIGRATION AND FATE OF INDICATOR CHEMICALS

Given the nature of contamination at ECC and the potential pathways of migration, indicator chemicals were assessed in terms of their behavior in soils, groundwater, and aquatic systems. Emphasis was placed on the mobility and persistence of each chemical. Mobility is important because it determines the rate of chemical migration away from the site. Persistence is important because it determines if a chemical will remain in the environment long enough to reach a receptor.

#### CHARACTERIZATION OF INDICATOR CHEMICALS

Table 5-2 lists some of the important physical-chemical properties of each indicator chemical. No inorganics were selected as indicators since only cadmium, lead, and zinc were found at concentrations above typical ranges in more than one sample. Considering the soils characteristic of the site and the physical-chemical properties of the inorganics, transport will be minimal.

Table 5-2
PHYSICAL-CHEMICAL PROPERTIES OF INDICATOR ORGANICS

		Boiling				
-	Molecular Weight	Point (°C) <sup>a</sup>	Vapor Pressure (torr)	Solubility (mg/L)	Log Kow <sup>C</sup>	Kd <sup>e</sup>
Volatile Organics						
1,1,2-trichloroethene	133.41	133.8	19 <mark>d</mark>	4,500 <sup>d</sup>	2.17	0.18
1,1,1-trichloroethane	133,41	74.1	97.0	480-4,400 <sup>d</sup>	2.17	0.18
Tetrachloroethene	165.83	121.0	14.ດີ	150-200	2.88	0.94
Trichloroethene	131.39	87.0	57.9 <sup>th</sup>	1,100	2.29	0.24
Toluene	92.13	110.6	28.7	535 <sup>f</sup>	2.69	0.60
Chloroform	119.38	61.7	150.5	8,200 <sup>f</sup>	1.97	0.12
Methylene chloride	84.99	39.8	350 0	20.000	1.25	0.022
Ethylbenzene	106.2	136.2	7 <sup>d</sup>	152 <sup>d</sup>	3.15	1.74
Acid Compounds						
Phenol	94.11	181.8	0.8 <sup>f</sup>	93,000 <sup>f</sup>	1.46	0.036
Base/Neutral Compounds						
Bis(2-ethylhexyl)phthalat	e 391.0	386.9	0.01 <sup>d</sup>	1.3 <sup>f</sup>	8.7	660,000
Dimethyl phthalate	194.2	282.0	0.01	90¢ <sup>£</sup>	2.12	0.16
Diethyl phthalate	222.2	298.0	0.05	4,320 <sup>f</sup>	3.22	2.05
Di-n-butyl phthalate	278.3	340.0	0.1 <sup>g</sup>	13	5.2	195
Other Organics						
PCB 1260	375.7	-	4.05x10 <sup>-5f</sup>	0.0027	7.14	17,000
1232	232.2	-	4.06x10 <sup>-3f</sup>	1.45	3.2	1.95

Boiling point at 760 torr.

btorr = 1 mm of mercury (Hg).

CKow = octanol-water partition coefficient.

dVapor pressure/solubility at 20°C.

eK = soil-water partition coefficient

fd
Vapor pressure/solubility at 25°C.

gVapor pressure/solubility @ 115°C

hVapor pressure/solubility @ 70°C.

GLT301/60

It is important to note that the actual migration and fate of the contaminants depend largely on the physical-chemical features of the site such as temperature, pH, percent soil moisture, geochemistry, soil type, and oxidation-reduction potential. Other factors that must be considered are potential reactions between chemicals and the formation of transformation byproducts. For example, under certain conditions tetrachloroethene is believed to breakdown to trichloroethene, and then to the "cis" form of dichloroethene and then to vinyl chloride. Each of the byproducts are compounds that would pose a health threat to receptors. It is beyond the scope of this project to research the migration and fate of all the byproducts; however, their significance should be recognized.

Table 5-3 provides a summary of the environmental behavior of the indicator organic compounds. Summaries are provided for three key sectors of the environment: subsurface soils and groundwater, surface soils, and aquatic systems. Potential transformation and transfer mechanisms are listed for each indicator chemical. Transformation mechanisms act to change the form of a chemical, while transfer mechanisms partition the chemical between media (e.g., volatilization is a water-air transfer; sorption is a water-soil transfer). The persistence of a chemical in a given sector of the environment is generally controlled by transformation mechanisms and volatilization. Chemical mobility in a given

#### Table 5-3 (Page 1 of 2) SUMMARY OF ENVIRONMENTAL BEHAVIOR OF INDICATOR ORGANIC COMPOUNDS IN SUBSURFACE SOILS, GROUNDWATER, SURFACE SOILS AND AQUATIC SYSTEMS

		Subsurface Soils and Groundwater			Surface Soils						
	Transformation			Transfer		Transformation				Transfer	
Compound	Oxidation	Hydrolysis	Biodegradation	Sorption	Oxidation	Hydrolysis	Photolysis	Biodegradation	Volatilization	Sorption	
1,1,1-Trichloroethane	I	6 toos - 1 yr	P <sup>a</sup>	I	I	P	ı	1	S	ı	
1,1,2-Trichloroethane	I	6 mos - 1 yr	P.ª	1	I	P	1	I	S	I	
Tetrachloroethene	8.8 mos	ı	P <sup>a</sup>	I	P	1	I	1	S	1	
Trichloroethene	10.7 mos	1	P <sup>a,</sup>	I	P	1	I	1	s	1	
Toluene	I	I	₽ <sup>b</sup>	1	P	I	P	P <sub>p</sub>	s	I	
Chloroform	I	1-3,500 yrs	Pª	I	I	P	1	p <sup>a</sup>	s	I	
Methylene Chloride	1	1-704 yrs	P	1	I	P	1	P	s	I	
Ethylbenzene	1	I	P	1	P	.I.	P	P	s	I	
Polychlorinated Biphenyls	1	I	days~mos	S	ı	1	$\mathbf{p}^{\mathbf{e}}$	days-mos	mos-yrs	s	
Phenol	I	1	s	I	P	1	P	S	P	I	
Phthalates	I	I	P	s <sup>f</sup>	I	P	I	P	I	s <sup>f</sup>	

Notes: S = Significant

I = Insignificant

M = Moderate

P = Possible

Under anaerobic conditions.
Under aerobic conditions.

GLT301/59-1

Clear, well aerated systems.

Waters high in iron and copper.

Depends on degree of chlorination.
Depends on the compound.

Table 5-3 (Page 2 of 2)

Aquatic S	y s	tez
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	Ndortt Discons								
		Tran	Transfer						
Compound	Oxidation	Hydrolysis	Photolysis	Biodegradation	Volatilization	Sorption			
1,1,1-Trichloroethane	I	6 mos - 1 yr	1	₽ <sup><b>a</b></sup>	min - hrs	I			
1,1,2-Trichloroethane	I	6 mos - 1 yr	I	P.ª	min - hrs	I			
Tetrachloroethene	8.8 mos	I	1	P.a.	min - hrs	1			
Trichloroethene	10.7 mos	1	I	P <sup>a</sup>	min - days	I			
Toluene	P	I	P ,	Pp	hrs	I			
Chloroform	1	1-3,500 yrs	I	P	min - hrs	I			
Methylene Chloride	I	1-704 yrs	1	P	min - hrs	I			
Ethylbenzene	P	I	P	P	5 - 6 hrs	I			
Polychlorinated Biphenyls	I	I	P <sup>e</sup>	days -mos b,e	mos - yrs	S			
Phenol	S	I	P	s	P	I			
Phthalates	I	P	1	P	I	sf			

Notes: S = Significant

I = Insignificant

M = Moderate

P = Possible

GLT301/59-2

Under anaerobic conditions.
Under aerobic conditions.
Clear, well aerated systems.
Waters high in iron and copper.

Depends on degree of chlorination,
Depends on the compound

sector is mainly controlled by sorption. Both tables list if the mechanism has a significant (S), insignificant (I), or moderate (M) impact on behavior. In cases where the significance is uncertain or dependent on environmental conditions, the mechanism is denoted as possible (P).

Environmental behavior profiles are provided in Appendix C for each indicator chemical. The following section summarizes site characteristics important to contaminant transport.

# KEY SITE CHARACTERISTICS

# Groundwater

The key site characteristics are rate of leachate flow to the shallow saturated zone and travel time of groundwater from the site to both the unnamed ditch and Finley Creek. Using an estimated 7.8 inches of recharge water per year to the shallow saturated zone, the leachate rate was calculated as 568 gallons per year per square foot (200 liters/per year per square meter). Groundwater velocities for the shallow saturated zone were calculated assuming flow from the eastern portion of the site is directed to the unnamed ditch and that flow from the northern and western portions is directed to either the unnamed ditch or Finley Creek. The average horizontal gradient for the eastern portion was

estimated to be 0.05 feet per foot and for the northwestern portion to be 0.02 feet per foot. An effective porosity of 0.20 was used and the average hydraulic conductivity was estimated as  $10^{-5}$  centimeters per second. The resulting groundwater velocities are 1.0 ft/yr for the northwestern portion and 2.6 ft/yr for the eastern portion. Contaminant velocities and travel times were then calculated using retardation factors.

In the shallow sand and gravel aquifer, the average hydraulic conductivity was estimated to range from  $10^{-2}$  to  $10^{-3}$  centimeters per second and the porosity was assumed to be 0.30. Using an average gradient of 0.03 feet per foot, the groundwater velocities were calculated to be around 100 to 1,000 feet per year.

# Surface Water

Flow rate in the unnamed ditch was estimated to be 0.1 cfs and flow in Finley Creek reportedly ranges from 0.1 to 4 cfs. Assuming a 1 fps velocity in the ditch, the travel time for contaminants to reach Finley Creek would be from 5 to 20 minutes. Finley Creek discharges into Big Eagle Creek which has a flowrate ranging from 25 cfs to 150 cfs. Big Eagle Creek eventually discharges into the Eagle Creek Reservoir which contains between 4.7 x 10 and 8.1 x 10 gallons of water.

## INDICATOR CHEMICAL TRANSPORT AND FATE

The site-specific behavior of the indicator chemicals can be discussed in terms of the profiles presented in Appendix C and some basic site characteristics. It is convenient to group the indicator chemicals as follows given similarities in their behavior: volatile organics, phenols, phthalates, and PCB's. Transport and fate of the indicator chemicals are based on a literature review and site characteristics. Due to the relatively limited literature available and the many estimates and assumptions necessary, the transport and fate calculated here are gross best estimates only. Actual transport and fate may vary by orders-of-magnitude.

# Methodology

In order to estimate degradation (whether biotic or abiotic) it was assumed that degradation occurred according to the formula:

$$C = C_{0} - ct$$

$$T_{m} + t$$

where: t = time

c = concentration at time t

c<sub>o</sub> = initial concentration

 $T_{m} = half-life; time when <math>c = \frac{1}{2}c_{o}$ 

Assuming the curve remains linear until  $t=T_m$ , half-life values were calculated from first order rates found in the literature (Half-lives were also obtained from the literature). Each indicator compound was researched individually and wherever possible degradation rates were obtained for each compound under the various site conditions (i.e., soil, groundwater, surface water, and air). When values could not be found in the literature, rates for those compounds were assumed to be the same as structurally similar compounds. Using average and maximum concentrations for  $c_0$  and the derived concentration for c, the above equation was solved for c. An example calculation is shown in Appendix c.

The methodology assumes that as a compound degrades and becomes limiting, the degradation rate approaches zero. The equation is an adaptation of the Michaelis-Merten relationship and results in a more conservative estimate of degradation than a simple first order rate equation. Each compound was considered the sole source in the degradation process and no effects from temperature changes or chemical interactions were considered.

# Volatile Organics

Although volatile organics were detected in ponding surface water, they should readily volatilize and should not persist in surface soils or ponding water at the ECC site.

Volatiles are present in elevated concentrations in the subsurface soil. Overall volatile organics exhibit high mobility and are therefore most likely to be leached out of the contaminated soil. If leaching is prevented, it is estimated that all the indicator volatiles except 1,1,1-TCA, TCE, and PCE will degrade (by either biodegradation or abiotic degradation) to acceptable levels within 10 years. (Acceptable levels are assumed to be concentrations in the soil which will not result in excess lifetime cancer risk greater than  $10^{-6}$  or daily chemical intakes greater than the acceptable daily intakes based on the assumptions outlined in the Endangerment Assessment in the next chapter.) Degradation of PCE to acceptable levels is estimated to take less than 10 years based on site average concentrations but from 100 to 500 years based on maximum concentrations. and 1,1,1-TCA are estimated to degrade to acceptable levels in less than 100 years and 20 years, respectively, at site average concentrations and from 500 to 1,000 years at maximum concentration.

Under existing site conditions, the volatile organics will leach from the unsaturated soil into the groundwater and slowly migrate towards the unnamed ditch or Finley Creek. Estimated concentrations in the groundwater are shown in Table 5-4. The travel times to reach the surface waters will vary greatly depending upon the compound, soil properties, the hydraulic conductivity, and the travel distance.

Table 5-4
ESTIMATED CONCENTRATIONS OF VOLATILE ORGANICS
IN GROUNDWATER DUE TO LEACHING

	Concentratio	n (ug/L)
Compound	Average	Maximum
G1 1 5	100	6 000
Chloroform	188	6,000
Methylene Chloride	101,250	3,500,000
1,1,1-TCA	40,250	1,100,000
1,1,2-TCA	25	825
TCE	100,000	275,000
PCE	4,000	50,000
Toluene	28,250	133,750
Ethylbenzene	5,500	38,000

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Estimates for travel time from the eastern portion of the site to the ditch are as follows:

methylene chloride	20 to 70 years
chloroform	30 to 120 years
1,1,1-TCA	40 to 150 years
1,1,2-TCA	40 to 150 years
TCE	50 to 200 years
toluene	100 to 400 years
PCE	150 to 600 years
ethylbenzene	300 to 1,000 years

These estimates were based on a hydraulic conductivity of  $10^{-5}$  cm/sec and would be an order-of-magnitude less using  $10^{-4}$  cm/sec (i.e., 2 to 7 years for methylene chloride). Estimates for travel time from the northwestern portion of the site to the ditch and Finley creek are as follows:

# o Unnamed Ditch:

methylene chloride	250	years
chloroform	400	years
1,1,1-TCA	500	years
1,1,2-TCA	500	years
TCE	650	years
toluene	1,300	years
PCE	2,000	years
ethylbenzene	3,400	years

# o Finley Creek:

methylene chloride	600	years
chloroform	1,000	years
1,1,1-TCA	1,300	years
1,1,2-TCA	1,300	years
TCE	1,600	years
toluene	3,200	years
PCE	5,000	years
ethylbenzene	8,400	years

Again these estimates were calculated using a hydraulic conductivity of  $10^{-5}\ \mathrm{cm/sec.}$ 

Considering the significant travel times, 1,1,1-TCA and 1,1,2-TCA will likely experience degradation given the magnitude of their hydrolysis rates. Because the groundwater is relatively shallow, it is likely that aerobic conditions exist. If this is the case, trichloroethene and tetrachloroethene will also experience some degradation. Degradation of the remaining volatiles does not appear likely, but will depend on the actual conditions. Some species of bacteria, for instance, have been shown to be able to degrade ethylbenzene.

Assuming no degradation of the compounds, concentrations in the surface waters following complete mixing with the groundwater are listed in Table 5-5.

Concentrations in the surface waters due to discharge of contaminated water from the shallow sand and gravel aquifer are estimated to be the following:

1,1-Dichloroethene	0.024	to	0.24 ug/L
TCE	0.064	to	0.64 ug/L
Methylene chloride	0.19	to	1.9 ug/L
PCE	0.03	to	0.3 ug/L

(assuming  $K = 10^{-3}$  cm/s)

Table 5-5
ESTIMATED CONCENTRATIONS OF VOLATILE ORGANICS IN SURFACE
WATER DUE TO GROUNDWATER DISCHARGE

Compound	Unnamed Ditch	Unnamed Ditch	Finley Creek <sup>C</sup>
Chloroform	0.31	0.1	0.01 to 0.1
Methylene Chloride	170	60	6 to 60
1,1,1-TCA	70	25	2.5 to 25
1,1,2-TCA	0.04	0.014	0.001 to 0.014
TCE	170	60	6 to 60
PCE	7	3	0.3 to 3
Toluene	50	16	1.6 to 16
Ethylbenzene	10	3	0.3 to 3

Concentrations in the unnamed ditch due to groundwater discharge from the eastern portion of the site.

NOTE: Concentrations were estimated assuming a hydraulic conductivity of 10<sup>-5</sup> cm/s. Values would be 10 times higher using 10<sup>-4</sup> cm/s.

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Concentrations in the unnamed ditch due to groundwater discharge from the northwestern portion of the site.

portion of the site.  $^{\rm c}$  Concentrations in Finley Creek due to groundwater discharge from the northwestern portion of the site. Concentrations vary depending on the flow rate.

The volatiles that reach the surface waters should readily volatilize if environmental conditions (e.g., temperature) are favorable. Any contaminants remaining in the surface water would experience a 1:8 to a 1:40 dilution upon entering Big Eagle Creek and then a further dilution within the Eagle Creek reservoir.

Once in the atmosphere, the volatiles should degrade via photooxidation. The volatiles should not be found in high concentrations in the surface water sediments. Figure 5-2 summarizes the transport and fate of the volatile organics at ECC.

# Phenols

Phenol in the subsurface soil is already below acceptable levels and is estimated to degrade at a rate which would result in only trace levels remaining after 5 to 10 years.

Phenol is easily desorbed from soil, however, and would readily leach into the groundwater. Phenol concentrations in groundwater are estimated to be about 80 mg/L based on average soil concentrations and about 4,000 mg/L based on maximum soil concentrations. Estimates of travel times from the eastern portion of the site to the unnamed ditch range from 20 to 80 years using a hydraulic conductivity of 10<sup>-5</sup> cm/sec. Travel time from the northwestern portion of the

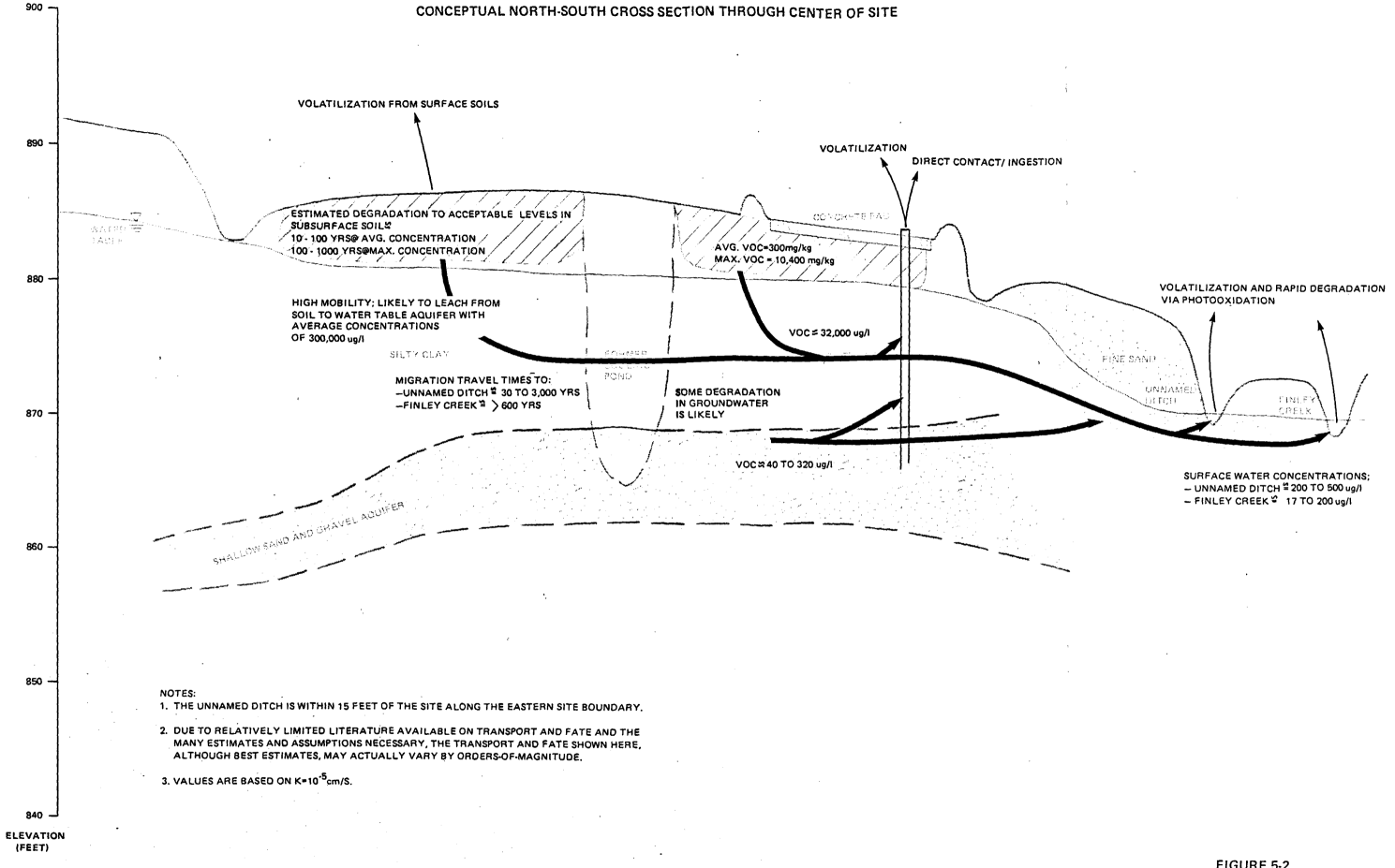


FIGURE 5-2 TRANSPORT AND FATE OF VOLATILE ORGANICS - ECC RI site to the ditch and Finley Creek using 10<sup>-5</sup> cm/sec are estimated to be 300 years and at least 650 years, respectively.

Degradation will most likely occur considering the long travel time and the biodegradability of phenol. Assuming no degradation, phenol concentrations are estimated to be on the order of 50 to 150 ug/L in the ditch. Phenol concentrations in Finley Creek following mixing are estimated to be about 5 to 50 ug/L. If degradation is considered, the concentrations are estimated to be in the range of 0.2 to 10 ug/L in the ditch, and 0.9 to 30 ug/L in Finley Creek depending on travel time ( $k = 10^{-5}$  cm/sec). Once in the surface waters phenol should degrade more rapidly since aeration enhances the reduction of phenol by microorganisms.

Volatilization of phenol will not be significant, nor will sorption in surface water sediments. Figure 5-3 summarizes the transport and fate of phenols at ECC.

#### Phthalates

Phthalate esters in the subsurface soil are already below acceptable levels and are estimated to biodegrade to trace levels within 100 to 200 years.

The phthalates found at ECC exhibit a range of physical-chemical properties. Bis(2-ethylhexyl) phthalate and di-n-butyl phthalate both have low solubilities and high soil-water partition coefficients. Diethyl and dimethyl phthalate have much higher solubilities and much lower partition coefficients. Consequently, the latter two exhibit some mobility within the environment and will leach from the contaminated soil into the groundwater. Only trace concentrations of bis(2-ethylhexyl)phthalate and di-n-butyl phthalate will appear in the groundwater:

	Concentration (ug/L)	
	Average	Maximum
Diethyl phthalate	50	1,100
Dimethyl phthalate	. 125	2,000
Di-n-butyl phthalate	0.50	10
Bis(2-ethylhexyl)phthalate	0.005	0.15

Estimated travel times from the eastern portion of the site to the unnamed ditch range from 40 to 150 years for dimethyl and 300 to 1,100 years for diethyl phthalate using  $10^{-5}$  cm/sec. Travel times for the northwestern portion of the site are orders of magnitude higher.

Degradation will most likely occur since biodegradation is a significant mechanism in the ultimate fate of the phthalate

esters. However, concentrations in the unnamed ditch are estimated to be less than 4 ug/L assuming no degradation. Estimated concentrations in Finley Creek are even lower and will be reduced considerably if degradation is considered.

Volatilization of phthalates will not be a significant pathway since they have very low vapor pressures. Phthalates should not be able to migrate to surface water sediments except in trace quantities unless there is direct runoff or discharge to the creek. Once in the surface water the phthalates will adsorb readily and tend to persist in the sediments. Figure 5-4 summarizes the transport and fate of phthalates at ECC.

# PCB's

PCB's will tend to persist in surface and subsurface soils. Some degradation may occur in onsite surface soils through volatilization, photolysis, and biodegradation. Subsurface degradation will be limited and (as with surface soils) will vary with the type of PCB mixture. Degradation to acceptable levels is estimated to take 50 years at site average concentrations and approximately 100 to 2,000 years at maximum concentrations.

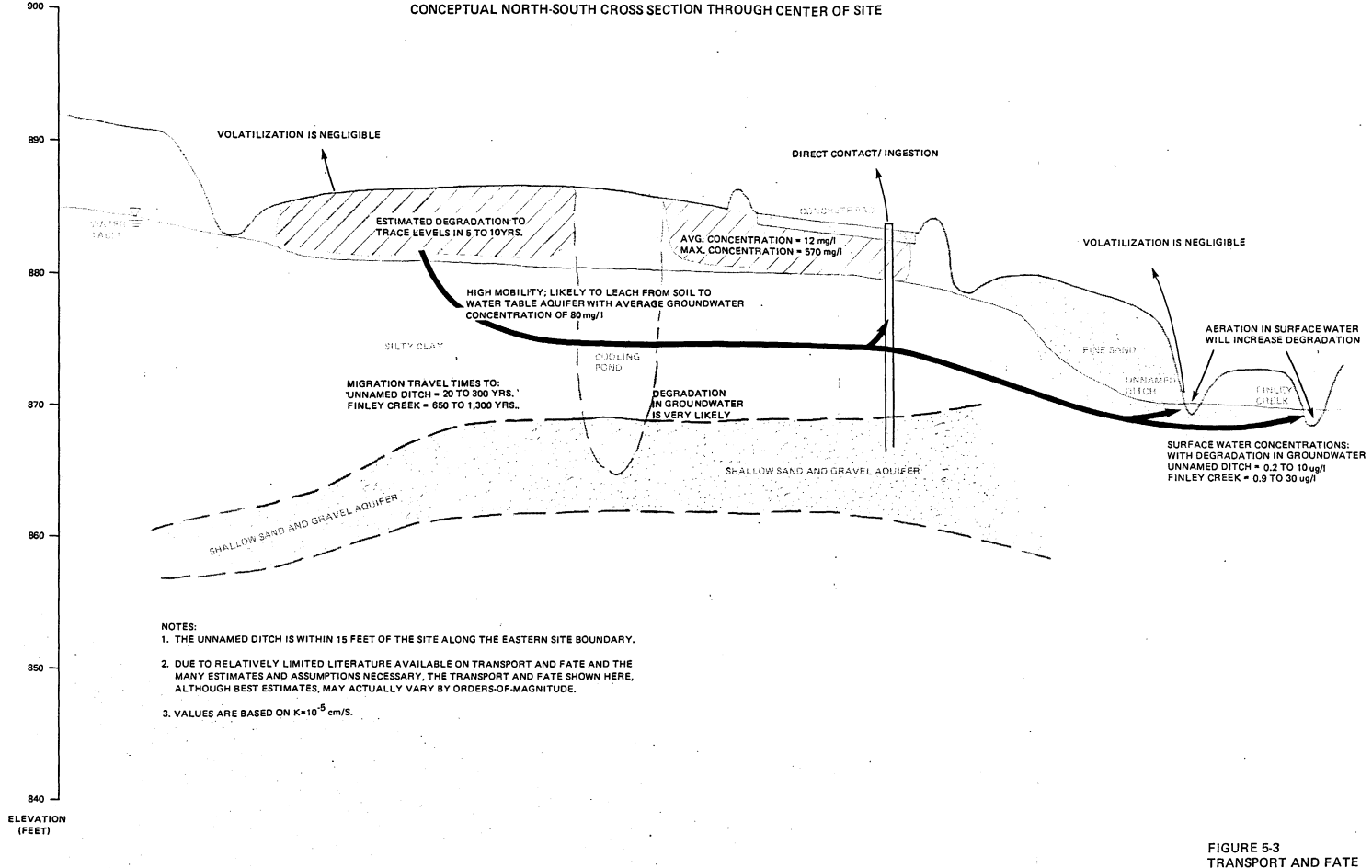


FIGURE 5-3
TRANSPORT AND FATI
OF PHENOLS
ECC RI

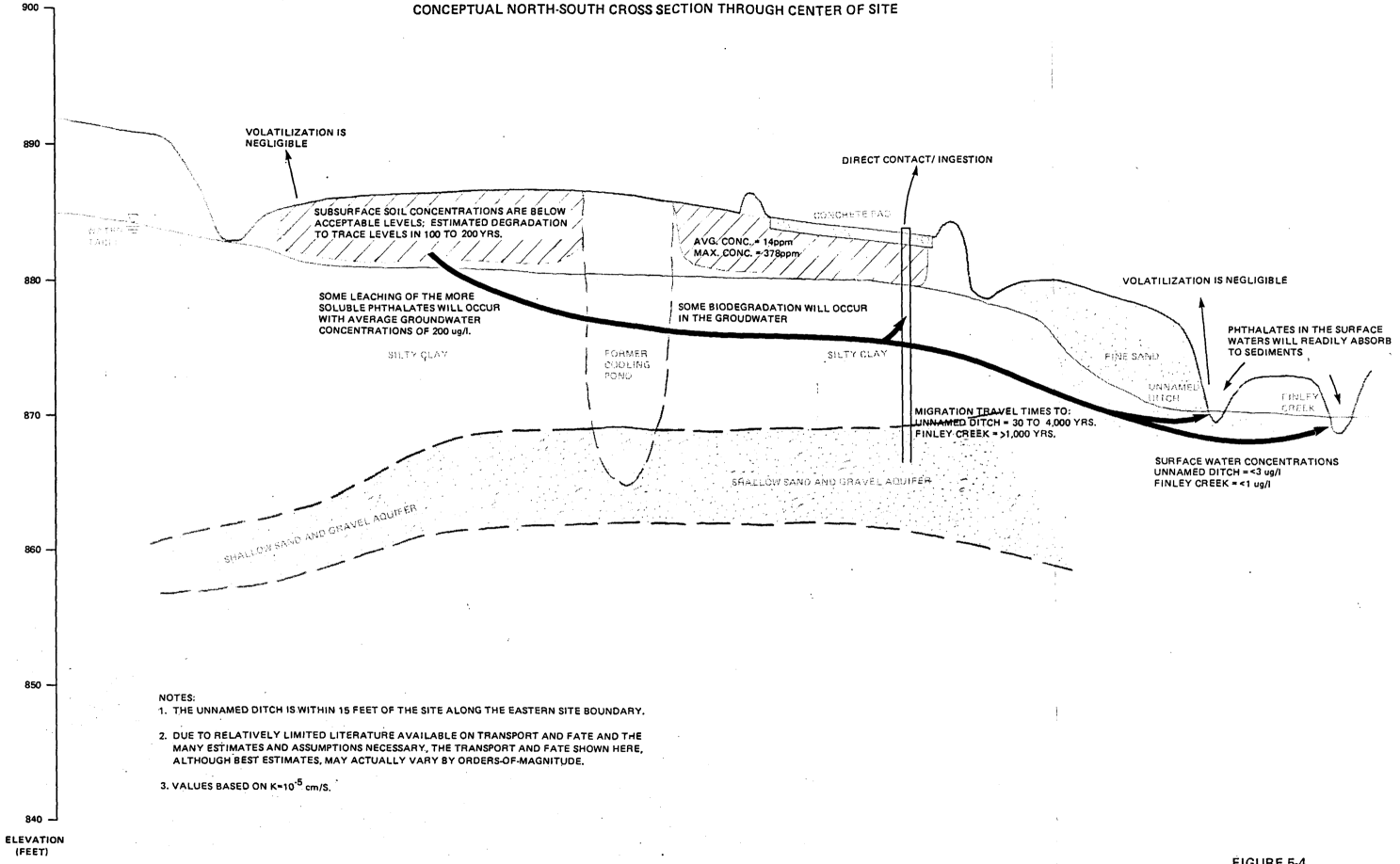


FIGURE 5-4
TRANSPORT AND FATE
OF PHTHALATES
ECC RI

PCB's readily adsorb to soil and have low solubilities. Of the two detected at ECC, only 1232 will leach into the groundwater and only in trace concentrations (25 ug/L based on average soil concentrations). PCB's are, however, not likely to migrate within the aquifer. If PCB's enter the ditch or creek by surface runoff or direct discharge, they would absorb readily to the sediments. Figure 5-5 summarizes the transport and fate of PCB's at ECC.

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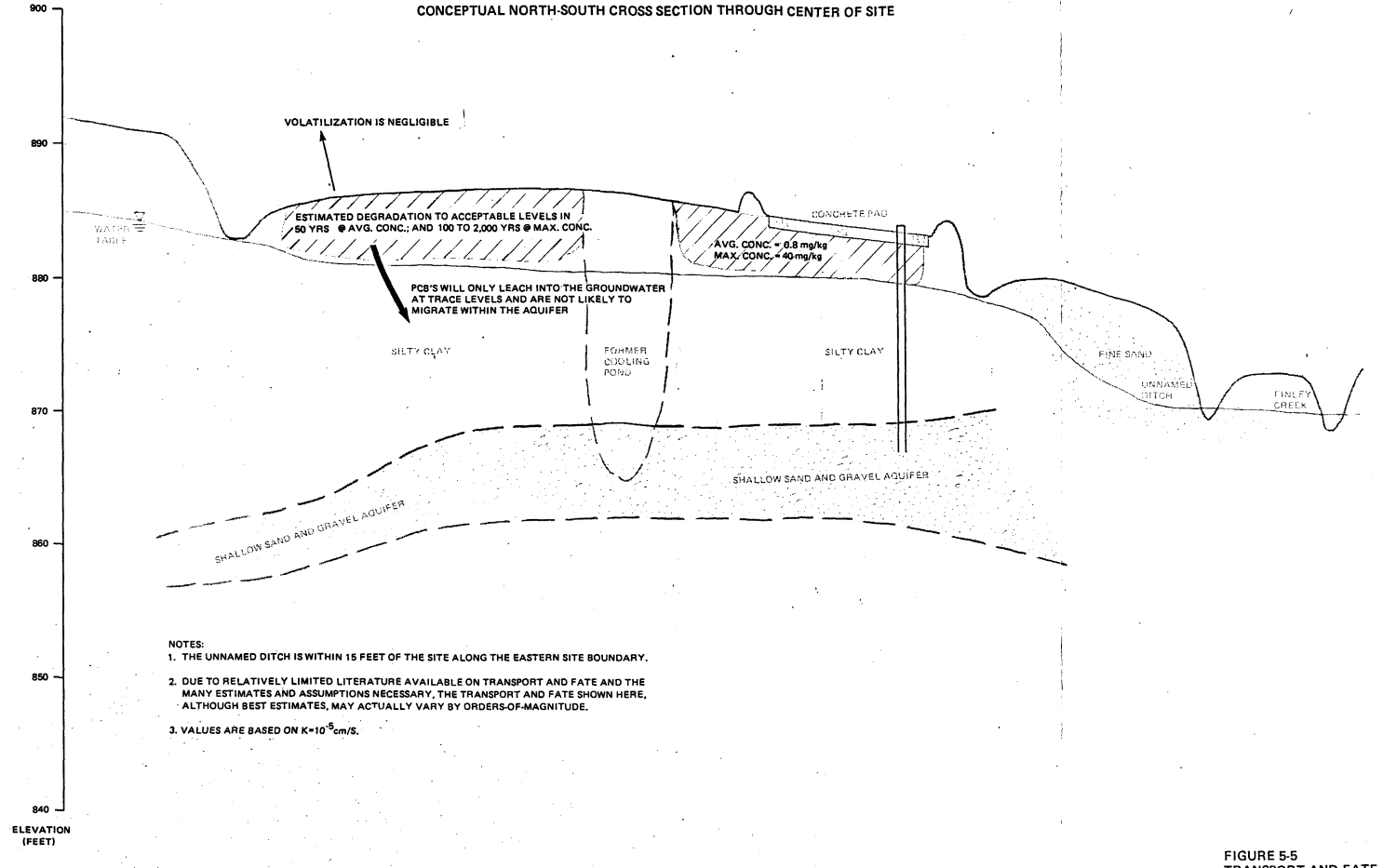


FIGURE 5-5
TRANSPORT AND FATE
OF PCB'S
ECC RI

#### Chapter 6

#### ENDANGERMENT ASSESSMENT

#### INTRODUCTION

This endangerment assessment analyzes the potential human health and environmental impacts of the ECC site in the absence of any remedial action (the no action alternative). It has two components, the public health evaluation and environmental assessment, which are discussed relative to each of the appropriate environmental media: soil, sediment, groundwater and surface water. Potential receptors are identified along with the hazardous substances present the environmental media. Both the quantitative and qualitative impact of contaminants on the public health and the environment are evaluated.

## **PURPOSE**

An endangerment assessment is a determination of the magnitude and probability of actual or potential harm to public health, welfare, or the environment by the threatened or actual release of a hazardous substance. Before taking action under Section 106 of CERCLA to abate the hazards or potential hazards at a site, the EPA must be able to properly document and justify its assertion that an imminent

hazard exists. The endangerment assessment provides this documentation and justification.

#### DEFINITION OF PROBLEM

Earlier chapters of this report have shown that environmental media at the ECC site have become contaminated with over 80 organic and inorganic chemicals (Table 6-1). The potential human health effects associated with exposure to many of these chemicals affect a range of human organ systems including the respiratory, nervous, circulatory, digestive, dermal, and urinary systems. Fourteen of the chemicals found at this site are potential human carcinogens (Table 6-2).

Chapter 5 of this report discusses the environmental fate and transport of site contaminants. The primary releases will be from soil to groundwater and then to surface water.

The population at risk consists of current and future human, plant, and wildlife populations residing on or adjacent to the ECC site. These populations are defined in greater detail in the public health evaluation and the environmental assessment in sections of this chapter.

Human exposure to contaminants is dependent on the environmental media in which the contaminant is present and the

Table 6-1 (Page 1 of 3)
SUBSTANCES DETECTED AT ECC DURING THE REMEDIAL INVESTIGATION

••	Compound	Soils	Sediments	Groundwater	Offsite Surface Waters
	VOLATILES	,			
Benzene				xs	
Chlorobe	enzene	X			
1,1,1-Tr	cichloroethane	X		S	S
1,1-Dich	loroethane	0		0	0
1,1,2-Tr	richloroethane	X			
Chloroet	hane			0	0
Chlorofo	orm	X		XS	
1,1-Dich	loroethene	. <b>X</b>		XS	
Trans-1,	2-Dichloroethene	0		S	s
	3-Dichloropropene	X		0	
Ethylben		X		XS	
-	ne Chloride	X		XS	
	cichloromethane		X		
	oroethene	X		XS	xs
Toluene		Х		XS	S
Trichlor	oethene	X		XS	XS
Vinyl ch	loride	X		XS	
Acetone		0		0	
	one (MEK)	0		os	
	-2-Pentanone	0		••	
Styrene				0	
o-Xylene	•	X		\$	0
2-Hexano		0		•	Ū
	-m-Cresol				
Phenol		x			
Benzoic	Acid	0	0		
2-Methyl		0	•		0
4-Methyl	-	0	0		0
BASE/NEU	TRALS				
1,2-Dich	lorobenzene	0			
Fluorant	hene			XS	
Isophoro	ne	X			

X = Substances quantitatively assessed for risk in endangerment assessment.

<sup>0 =</sup> Substances not quantitatively assessed because a cancer potency or acceptable daily intake value is not available.

S = Substance compared to standard, criteria, or guideline.

Table 6-1 (Page 2 of 3)

				Offsite
Compound	Soils	Sediments	Groundwater	Surface Waters
	<del></del>	<del></del> _		
Naphthalene	0		XS	
bis(2-Ethylhexyl) Phthalate	X	X	XS	
Benzyl Butyl Phthalate	0			
di-n-Buyl Phthalate	X			
di-n-Octyl Phthalate	0			0
Diethyl Phthalate	X		XS	
Dimethyl Phthalate	X			
Crysene	0		S	
Benzo(ghi) Perylene	0			
Fluorene	0			
Phenanthrene	0			
Pyrene			S	
2-Methylnapththalene	0			
PCB'S/PESTICIDES				
PCB-1232	x			
PCB-1260	X			
INORGANICS				
Antimony	x	x	XS	
Arsenic	X	X	S	
Aluminum	0	0	0	0
Barium	0	0	S	
Berylium	Х	X		
Cadmium	Х	X		
Cobalt	0	0	0	
Calcium	0		0	
Chromium	X	X	XS	
Copper	0		S	
Iron	0	0	S	0
Lead	X	X	XS	
Magnesium			0	

X = Substances quantitatively assessed for risk in endangerment assessment.

<sup>0 =</sup> Substances not quantitatively assessed because a cancer potency or acceptable daily intake value is not available.

S = Substance compared to standard, criteria, or guideline.

Table 6-1 (Page 3 of 3)

Compound	Soils	Sediments	Groundwa	Offsite Surface ter Waters
• •				
Manganese	0	0	S	0
Potassium				
Sodium			0	
Nickel	Х	X	XS	
Selenium		X	XS	
Mercury	Х	X	XS	
Silver		Х	X	
Thallium		X	XS	
Tin		0		
Vanadium	0	0		
Zinc		X	S	
Cyanide	х	X		

X = Substances quantitatively assessed for risk in endangerment assessment.

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<sup>0 =</sup> Substances not quantitatively assessed because a cancer potency or acceptable daily intake value is not available.

S = Substance compared to standard, criteria, or guideline.

Table 6-2 POTENTIAL CARCINOGENS DETECTED AT ECC

	Carcinogen By U.S. EPA Carcinogen Assessment Group	fo	r Rese	nal Age arch or ategory 2B	h
Benzene	X	Х	X	X	Х
1,1,2-Trichloroethane	X				X
Chloroform	X			X	
1,1-Dichloroethene	X				X
Methylene Chloride	X				X
Tetrachloroethene	X				Х
Trichloroethene	X				X
Vinyl chloride	X	X			
PCB (Total)	X			X	
Arsenic _	X	X			
Berylium <sup>C</sup>	X		X		
Cadmium <sup>C</sup> _	X		X		
Chromium <sup>C</sup>	X	X			
Nickel <sup>C</sup>	X			X	

<sup>&</sup>lt;sup>a</sup>Potencies set by U.S. EPA Carcinogen Assessment Group (CAG) (U.S. EPA, Dec. 1984).
DInternational Agency for Research on Cancer Classification

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<sup>(</sup>WHO 1982):

<sup>1 -</sup> Human carcinogen

<sup>2</sup>A - Probable human carcinogen, positive animal carcinogen with limited evidence of human carcinogenicity.

<sup>2</sup>B - Probable human carcinogen, positive animal carcinogen with insufficient data on human carcinogencity.

<sup>3 -</sup> Data inadequate to be classified as to carcinogenicity in humans.

Carcinogen by inhalation route only.

current and future use of the site and adjacent property.

Contact with contaminants by natural population is governed by the environmental media contacted and the habitat and range of the population. The potential exposure pathways at ECC are listed in Table 6-3.

#### PUBLIC HEALTH EVALUATION

#### INTRODUCTION

The intent of the public health evaluation is to identify potential threats to human health in the absence of remedial action. It is assumed that the site has the potential for unrestricted future development under the no action alternative. This public health evaluation section characterizes the population at risk, describes the risk assessment approach, and presents summaries of the public health risk by media. Appendix E presents the risk assessment by media in greater detail.

#### Population at Risk

The ECC site is in Union Township of Boone County, Indiana. The 1982 population of Union Township was 1,827. There are no population projections available for Union Township at present, however, based on past trends the population of

#### Table 6-3 (Page 1 of 2) POTENTIAL EXPOSURE PATHWAYS

Release Source	Transport Media	Exposure Point	Exposure Route	Potential Population Exposed
1 Fugitive Dust	Air	Onsite and Offsite	Inhalation	Human-current and future
• •	Air	Onsite and Offsite	Ingestion	Human-current and future
2 Volatilization from soil	Air	Onsite and Offsite	Inhalation	Human-current and future
3 Site runoff	Surface Water	Unnamed ditch/ Finley Creek/ Eagle Creek	Direct contact (dermal absorption)	Human-current and future
4 Site runoff	Surface Water	Unnamed ditch/ Finley Creek/ Eagle Creek	Inhalation of volatilize compounds (intermedia transfer to air)	Human-current and future
5 Site runoff	Surface Water (fish)	Unnamed ditch/ Finley Creek/ Eagle Creek	Ingestion of fish	Human-current and future
6 Site runoff	Surface Water	Unnamed ditch/ Finley Creek/ Eagle Creek	Direct contact/ ingestion	Fish and other aquatic species
7 Soil	Direct content	Onsite	Dermal absorption	Human-current and future
8 Soil	Direct content	Onsite	Ingestion	Human-current and future
9 Soil	Direct content	Onsite	Ingestion	Terrestrial species
10 Groundwater	Discharge to surface water	Unnamed ditch/ Finley Creek/ Eagle Creek	Same as #3, 4, 5, 6	
11 Groundwater	Direct Use (wells)	Onsite	Ingestion	Human-current and future
12 Groundwater	Direct Use (wells)	Onsite	Inhalation	Human-current and future
13 Groundwater	Direct Use (wells)	Onsite	Dermal Absorption	Human-current and future
14 Groundwater	Direct Use (wells)	Offsite	Same as #11, 12, 13	Human-current and future
GLT/360/66				

Union Township could double by the year 2000. The zoning for the area around the site is shown in Figure 6-1.

There are approximately 30 residences within a 1/2 mile radius of the ECC site. Assuming development of 1 acre lots, the number of residences within a 1/2 mile radius of the ECC site could increase to around 300. There are currently no hospitals, schools, or nursing homes in the immediate area. Residents could become potentially at risk if they contacted contaminated soil, groundwater, surface water or biota on or adjacent to the ECC site. Exposure will be limited by location of residence (example: upgradient versus down gradient from site), lifestyle (example: fishing versus not eating fish), and frequency of contact.

The unnamed ditch flows into Finley Creek which empties into Big Eagle Creek. Big Eagle Creek ultimately flows into Big Eagle Creek Reservoir which is one of the drinking water sources for Indianapolis. If contaminants reach the reservoir then users of the reservoir could be at risk.

#### Approach

The concentration of contaminants found in the environmental media during the remedial investigation as well as concentrations of contaminants projected (see Chapter 5) for

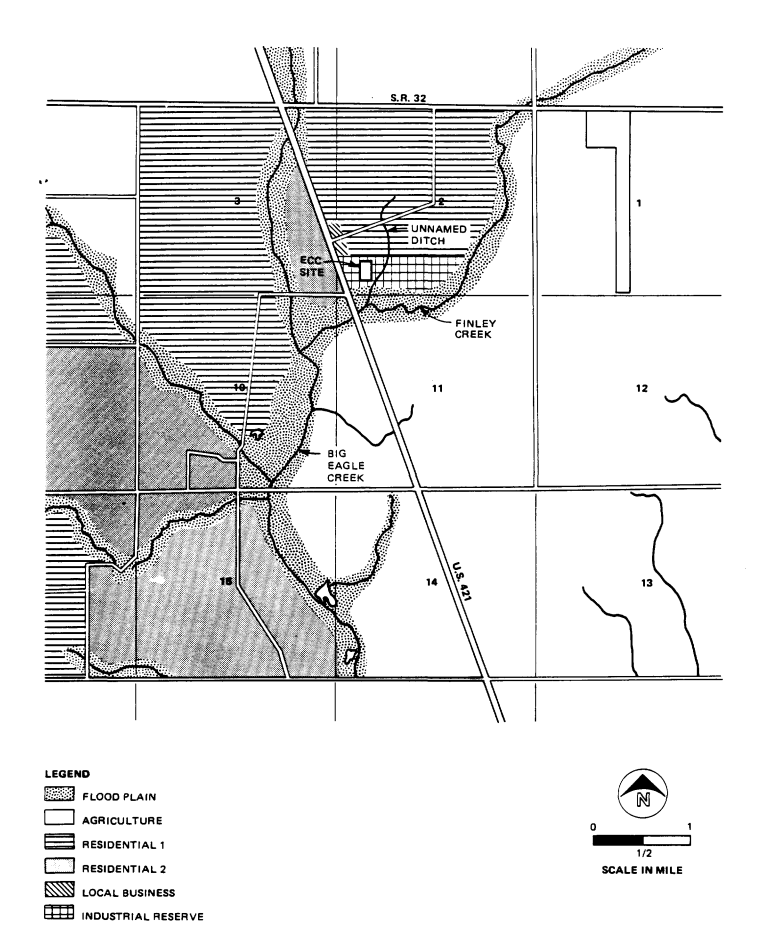


FIGURE 6-1 ZONING ADJACENT TO ECC IN UNION TOWNSHIP ECC RI

those media on the basis of the environmental fate and transport are used in this evaluation. Complete exposure routes are assessed using both present and predicted concentrations of contaminants at exposure points.

The concentration of chemicals at exposure points is compared to relevant or applicable standards, criteria, and guidelines where appropriate. These include the Safe Drinking Act Maximum Contaminant Levels (MCL's), and Clean Water Act Ambient Water Quality Criteria.

The exposed population's current and projected intake of selected compounds is estimated. This is performed for carcinogenic compounds and toxicants (noncarcinogens).

For the carcinogens present that are given cancer potencies by the U.S. EPA Carcinogen Assessment Group (CAG) (U.S. EPA, December 1984), an excess lifetime cancer risk is calculated by each appropriate exposure route. Excess lifetime cancer risk is defined as the incremental increase in the probability of getting cancer compared to the probability if no exposure occurred. For example, a 10<sup>-6</sup> excess lifetime cancer risk would represent the risk resulting from an exposure that would increase cancer incidence by one case per million people exposed. The equation used for the estimation of excess lifetime cancer risk assessments is:

Risk = 1 - exp(-[dose x cancer potency])

The use of this equation for computing risk is presented in Appendix D.

A comparison is made, by exposure route, between the projected intakes for the potentially exposed population and the acceptable intakes for each toxicant (noncarcinogen) for which an acceptable daily intake (ADI) has been established. An ADI is the amount of toxicant (in mg/day for a 70-kg person) that is not anticipated to result in any adverse effects after chronic exposure to the general population including sensitive subgroups (Dourson, Stara, 1983).

Some compounds do not have ADI's, cancer potencies, or standards and criteria. Of these compounds, those which are in significant concentrations or are of toxicological/public health importance are examined qualitatively.

Two exposure settings are defined to estimate the potential risks from development and use of the site and the areas adjacent. The residential setting assumes the potential for construction of residences at or adjacent to the site. This includes excavation of contaminated subsoil which could be placed into a garden or child play area. Residents could inadvertently ingest contaminated soil during outside activities and soil could be transported into the home on hands,

clothing, or by pets. Exposure to soil, however, is limited by weather conditions. It is assumed that the shallow groundwater below the site is used for domestic wells.

The adult worker setting assumes that a light industrial or commercial development occurs at the site. As in the residential setting, subsurface soil may be excavated during the construction and left on the surface and the shallow groundwater is used for wells. The workers are assumed to spend a significant part of their day in outdoor activities, but their exposure to the soil is also limited by weather as well as duration of work periods.

#### Limitations

When assessing public health risk it is reasonable to be conservative and assess upper bound situations. The risk assessment process uses specific assumptions, generalizations and recognized standard estimations. These assumptions and estimations are listed in Table 6-4.

The risk assessment process involves considerable uncertainty. The uncertainty is derived from availability of data, scientific judgments and assumptions that may or may not accurately reflect actual conditions. A listing of these uncertainty factors is presented in Appendix D.

### Table 6-4 (Page 1 of 2) RISK ASSESSMENT ESTIMATION AND ASSUMPTIONS

#### Assumption or Estimation

#### Comment

Exposure constant over 70 years

Concentration of contaminants constant over 70 years

Absorbed dose equal to 100% of amount ingested

Years in lifetime = 70 Adult body weight = 70 kg Adult water consumption -2L/day

Soil consumption:
10 grams/day/ - "pica" child;
1 gram/day/average child;
0.1 gram/day/adult;
0.5 gram/day/adult worker.

For carcinogens: lifetime average water ingestion rate (LAWI)=0.035 L/kg-body weight/day; lifetime average soil ingestion rate (LASI)=0.028 g/kg body weight/day

Correction of LASI to account for climatic influence: 0.013 g/kg body weight/day for residential setting; 0.00013 g/kg body weight/day for occupational setting.

In calculating downstream concentrations of contaminant dilution is only mechanism for reducing concentration. Conservative assumption.

Conservative assumption. Not all degradation rates are available.

Values for absorption efficiency are not readily obtainable. Using absorption efficiency as low as 25 percent would not reduce the excess lifetime cancer risk level by an order-of-magnitude.

U.S. EPA standard values used in deriving risk

Based on work of Kimbrough, et. al. (1984), and Schaum (U.S. EPA, 1983).

These are age and time weighted rates over a 70 year lifetime to account for the relatively higher ingestion rates per kg of body weight in younger age classes (see Appendix D).

See Appendix D.

Conservative assumption.
Actually volatilization would
be the major environmental
fate of volatile compounds.

#### Assumption or Estimation

#### Comment

The site has the potential for unrestricted future residential and commercial development.

Part of the definition of no action.

In assessing projected release of contaminants from soil to groundwater, the contaminants are treated as if they release at the same rate. The actual release ratio vary by contaminant due to physical and chemical characteristics.

No degradation in groundwater.

Conservative assumption to assess upper bound risk.

Contaminants release at the same rate from groundwater to surface water.

The actual release rates vary due to physical and chemical characteristics. Assumption made to keep assessment simple.

Maximum concentrations in groundwater are resprentative of entire zone.

Conservative assumption to assess upper bound risk.

Values of less than quantification limit are treated as if they are equal to the quantification limits. Conservative assumption to assess upper bound risk.

Dilution of groundwater to unnamed ditch is 1:600.

Based on estimated groundwater flow and estimated flow in the unnamed ditch.

Dilution of unnamed ditch to Finley Creek ranges from 1:2 to 1:40 and the dilution of Finley Creek to Eagle Creek ranges from 1:40 to 1:130. Based on limited USGS stream gaging.

No volatilization of compounds in surface water.

Conservative assumption to assess upper bound risk. Volatilization is likely.

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#### SOIL

The soil assessment is limited to subsurface soil exposure. Exposure to contaminated subsurface soils could only occur if the site is developed and soils are excavated. There is indirect evidence from the site surface water data that the "clean cover" of imported material in the northern area of the site may be contaminated. Without soil data this surface material cannot be assessed.

The ECC site was separated into two areas, northern (covered by imported material) and southern (covered by cement pad) (see Figure 4-2), for the evaluation of potential exposure of the public to site contaminants in the subsurface soils. The analysis is based on average and maximum contaminant concentrations found in the soil test pits in the northern area and the soil borings in the southern area.

For assessing the exposure to contaminated soil, the residential lifetime soil ingestion rate is estimated as 0.013 g/kg body weight/day (about 9 ounces per year) and the occupational lifetime soil ingestion rate is estimated as 0.00013 g/kg body weight/day (about one-tenth ounce per year). Adult soil ingestion could be as low as zero. It is assumed that exposure to contaminated soil is limited by climatic conditions such as precipitation, or frozen ground. In this area of Indiana, conditions suitable to limit

exposure occur 53 percent of the year (NOAA, 1980). See
Table 6-4 for exposure assumptions and Appendix D for a more
detailed description of derivation of soil exposures.

#### Ingestion

If the site is developed, outdoor activity on or adjacent to the site by people and pets provides access to contaminated soils. Contaminanted soil may be airborne during dry periods and adhere to hands and clothing, or it can be inhaled and inadvertently ingested.

A summary of the estimated risks attributed to ingestion of contaminated soil is shown in Table 6-5 (see Appendixes D and E for more detail on the derivation of risks). For example, the excess lifetime cancer risk for a residential setting from the soils in the northern portion of the ECC site could be  $4 \times 10^{-3}$  for maximum concentrations and  $4 \times 10^{-4}$  for average concentrations. The primary chemicals contributing to the risk are tetrachloroethene, trichloroethene, and PCB's.

Estimated daily chemical intakes in Table 6-6 show that xylenes, lead, and ethylbenzene exceed published Acceptable Daily Intakes (ADI's) at the ingestion rate of 10 grams of soil per day and xylenes and lead exceed ADI's at the 1 gram per day ingestion rate.

Table 6-5 (Page 1 of 2)
SUMMARY OF EXCESS LIFETIME CANCER RISK FROM INGESTION OF SOIL
FROM THE ECC SITE

Contaminant Concentration Scenario	Setting	Location	Major Chemicals of Concern	Total Excess Lifetime Cancer Risk
Maximum	Residential	Southern Area Intermediate Soil Depth	Chloroform Tetrachloroethene Trichloroethene	4 x 10 <sup>-5</sup>
Maximum	Occupational	Southern Area Intermediate Soil Depth	Chloroform Tetrachloroethene Trichloroethene	4 x 10 <sup>-7</sup>
Average	Residential	Southern Area Intermediate Soil Depth	Chloroform Tetrachloroethene Trichloroethene	8 x 10 <sup>-6</sup>
Average	Occupational	Southern Area Intermediate Soil Depth	Chloroform Tetrachloroethene Trichloroethene	8 x 10 <sup>-8</sup>
Maximum	Residential	Southern Area Deep Soil Depth	Trichloroethene Chloroform Tetrachloroethene	3 x 10 <sup>-8</sup>
Maximum	Occupational	Southern Area Deep Soil Depth	Trichloroethene Chloroform Tetrachloroethene	3 x 10 <sup>-10</sup>
Average	Residential	Southern Area Deep Soil Depth	Trichloroethene Chloroform Tetrachloroethene	6 x 10 <sup>-9</sup>
Average	Occupational	Southern Area Deep Soil Depth	Trichloroethene Chloroform Tetrachloroethene	6 x 10 <sup>-11</sup>

Table 6-5 (Page 2 of 2)

Contaminant Concentration Scenario	Setting	Location	Major Chemicals of Concern	Total Excess Lifetime Cancer Risk
Maximum	Residential	Northern Area Shallow Soil Depth	PCB Trichloroethene Tetrachloroethene	4 x 10 <sup>-3</sup>
Maximum	Occupational	Northern Area Shallow Soil Depth	PCB Trichloroethene Tetrachloroethene	4 x 10 <sup>-5</sup>
Average	Residential	Northern Area Shallow Soil Depth	PCB Trichloroethene Tetrachloroethene	3 x 10 <sup>-4</sup>
Average	Occupational	Northern Area Shallow Soil Depth	PCB Trichloroethene Tetrachloroethene	3 x 10 <sup>-6</sup>
Maximum	Residential	Northern Area Intermediate Soil Depth	PCB's Arsenic	8 x 10 <sup>-4</sup>
Maximum	Occupational	Northern Area Intermediate Soil Depth	PCB's Arsenic	8 x 10 <sup>-6</sup>
Average	Residential	Northern Area Intermediate Soil Depth	PCB's Arsenic	2 x 10 <sup>-5</sup>
Average	Occupational	Northern Area Intermediate Soil Depth	PCB's Arsenic	2 x 10 <sup>-7</sup>

Table 6-6
SUMMARY OF COMPOUNDS EXCEEDING ACCEPTABLE DAILY INTAKE
FROM SOIL INGESTION AT THE ECC SITE

				Daily Chemical Intakes Using Maximum Concentrations					rage Chemical Int Average Concentr	
Location	Chemical	ADI (ug/day)	Maximum Concentration ug/kg	@ 10 gm Soil/Day (ug/day)	@ 1 gm Soil/Day (ug/day)	@ 0.1 gm Soil/Day (ug/day)	Minimum Concentration ug/kg	@ 10 gm Soil/Day (ug/day)	@ 1 gm Soil/Day (ug/day)	@ 0.1 gm Soil/Day (ug/day)
South Pad Intermediate Depth							-			
South Pad Deep Depth	•		-				-			
North Test Pits										
Shallow Depth	Ethylbenzene	9,500	1,500,000	15,000	1,500	150	145,800	1,458	149	15
	Xylenes	1,200	6,800,000	68,000	6,800	680	629,900	6,299	630	63
	Lead	100	376,200	3,762	376	38	71,700	717	72	7
North Test Pits										
Intermediate Depth	Cadmium	170	27,000	270	27	3	3,900	39	4	0.4
•	Lead	100	415,200	4,152	415	41	60,200	602	6	6

Compounds present did not exceed ADI.

•

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#### Dermal Absorption

The amount of soil that comes in contact with human skin depends on factors such as behavior, soil type, weather conditions, and exposed skin area. These factors are highly variable, therefore estimation of dermal soil contact is difficult. Additionally, dry absorption rates for the variety of compounds found in the soil are not available. The data that do exist are based primarily on animal studies and extrapolated to humans which introduces uncertainty because of differences in skin properties. Because of these factors, a quantification of risk associated with dry absorption of compounds in soil is impractical. Only the qualitative statement that dermal exposure could increase risk can be made.

#### Dust Inhalation

Variables such as wind erosion, the organic content of soil, exposed surface area, and body absorption mechanisms make quantification of risk from dust inhalation difficult and tenuous. Only the qualitative statement that exposure through dust inhalation could increase risk can be made.

#### SEDIMENT

For the assessment of human exposure to sediment, the unnamed ditch and the Finley Creek sampling points are treated as separate exposure points. The analysis is based on maximum sediment contaminant concentrations attributable to the ECC site at each point. The maximum concentrations are used due to the limited number of sample points. It is assumed that residences and work places are or could be adjacent to areas of contaminated sediment and sediment may not be covered by water during low flow periods of the year.

As with soils, both residents and adult workers in the area, could incur health risks resulting from exposure to contaminated sediment during outside activities, or if sediment is transported into houses on hands, clothing, or by pets. The ingestion rates developed for soils are also used for sediments.

#### Ingestion

As a result of outdoor activity adjacent to the streams and river, people and pets have access to contaminated sediment. Contaminated sediment may be airborne during dry periods and adhere to hands and clothing or be ingested.

A summary of the estimated risks attributed to ingestion of contaminated sediments is shown in Table 6-7, (see Appendixes D and E for more detail on the derivation of the risks.) For example, the excess lifetime cancer risk for the residential setting near sampling point 004 in Finley Creek is than  $2 \times 10^{-11}$  for maximum concentrations. The primary chemical contributing to the risk is methylene chloride. Estimated daily chemical intakes in Table 6-8 show that lead exceeds a derived ADI at sampling point 004.

#### Dermal Absorption and Dust Inhalation

The same restrictions on the quantification of risk for the dermal absorption and inhalation of soils also is true for sediments.

#### GROUNDWATER

Groundwater is a major transport and release media for contaminants from the ECC site. The shallow saturated zone and the shallow sand gravel aquifer are the two portions of the groundwater impacted by contaminants from the ECC site.

Over 40 compounds are found in the groundwater with the volatile compounds being of most concern. Any risk from groundwater comes from it's direct use or the discharge of groundwater to surface waters. Direct use of groundwater

Table 6-7
SUMMARY OF EXCESS LIFETIME CANCER RISK
FROM EXPOSURE TO SEDIMENT FROM ECC SITE

Contaminant Concentration Scenario	Setting	Location	Major Chemicals of Concern	Total Excess Lifetime Cancer Risk
Maximum	Residential	003	Methylene Chloride	5 x 10 <sup>-4</sup>
Maximum	Occupational	003	Methylene Chloride	5 x 10 <sup>-3</sup>
Maximum	Residential	004	Methylene Chloride	2 x 10 <sup>-11</sup>
Maximum	Occupational	004	Methylene Chloride	2 x 10 <sup>-13</sup>
Maximum	Residential	005	Methylene Chloride	7 x 10 <sup>-11</sup>
Maximum	Occupational	005	Methylene Chloride	7 x 10 <sup>-13</sup>

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Table 6-8
SUMMARY OF COMPOUNDS EXCEEDING ACCEPTABLE DAILY INTAKE
FROM SEDIMENT INGESTION AT THE ECC SITE

Daily Chemical Intakes

				Using Max	cimum Observed Concen	trations
			Maximum	@ 10 gm	@ 1 gm	@ 0.1 gm
		ADI	Concentration	Soil/Day	Soil/Day	Soil/Day
Location	Chemical	(ug/day)	ug/kg	(ug/day)	(ug/day)	(ug/day)
SD003	a		-			
SD004	Lead	100	15,500	155	. 15	1
SD005	a		-			

a Compounds present did not esceed ADI.

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would include either consumption for drinking and cooking or from bathing. The discharge of groundwater to surface water is addressed in the surface water section.

The current population at risk are the users of one domestic well down gradient from the site, but prior to the discharge of the aquifer to Finley Creek. Domestic well sample results do not show any evidence of contaminants reaching this well. It would appear that this exposure pathway is currently incomplete.

Because of this, only future groundwater use, either residential or occupational, is considered. The size of the population that could use the groundwater would be limited by the relatively small area underlain by the aquifer between the ECC site and the aquifers discharge to surface water.

Risks are based on current data from the RI and projected release of contaminants from the soil to the groundwater as estimated in Chapter 5. Well Nos. 8A, 9A, and 10A represent the shallow sand and gravel aquifer and well No. 11A represents the shallow saturated zone (see Appendix E). For both zones, contaminant concentrations found during the RI in these wells are used to estimate risk under current conditions. The projected releases to the shallow saturated zone are used to estimate risk under future conditions in that

aquifer. The maximum concentrations are used from the RI data and maximum and average concentrations are used for the projected releases to the groundwater.

Appendix D presents derivation of ingestion and dermal absorption exposures. Appendix E presents the risk assessment for the groundwater in detail. A summary is presented below.

#### Comparison to Standards, Criteria, and Guidelines

Table 6-9 compares the maximum value for each compound found in wells representing the onsite aquifers (both shallow saturated zone and shallow sand and gravel) to relevant or applicable standards, criteria, and guidelines for the consumption of water.

Iron exceeds the secondary MCL, which is not a health based standard. This level is also found in the upgradient wells and represent areawide concentrations. 1,2-trans dichloroethene and trichloroethene exceed the proposed MCL's. Methylene chloride and trichloroethene exceed the AWQC  $10^{-6}$  cancer risk level. Trichloroethene also exceeds the chronic health advisory level.

Table 6-9
COMPARISON OF GROUNDWATER TO STANDARDS AND CRITERIA

Compound	Maximum Concentration	SDWA <sup>b</sup> MCL Primary	SDWA <sup>C</sup> MCL Secondary	AWQC <sup>d</sup> Toxicity	AWQC <sup>e</sup>	Health f Advisory	Criteria Exceeded
SHALLOW SAND AND GRAVEL AQUIFER:							
Barium	353	1,000	-	-	-	-	Y
Chromium	13	50	-	50	-	-	N
Iron	2,545	-	300	-	-	-	Y
Manganese	40	-	50	-	-	-	N
Nickel	46	-(-)	_	15.4	-	-,	Y
1,1-dichloroethene	8	7 <sup>(h)</sup>	_	-	0.033	70(j)	Y
Methylene chloride	64	-	-	-	0.19	150 <sup>(j)</sup>	Y
Tetrachloroethene	9	-(-)	•	-	0.8	20(j)	Y
Trichloroethene	21	5 <sup>(h)</sup>	-	-	2.8	75 <sup>(j)</sup>	Y
SHALLOW SATURATED ZONE - CURRENT CONCENTRATION	s:	<i>(</i> 1.)				4.5	
Trichloroethene	28,000	5 <sup>(h)</sup>	-	-	2.8	75 <sup>(j)</sup>	Y
SHALLOW SATURATED ZONE - PROJECT CONCENTRATION	S:						
Chloroform	6,000(188)	100 <sup>(i)</sup>	-	-	0.19	-,	Y
Methylene chloride	3,500,000(101,250)		-	-	0.19	150(j)	Y
1,1,1-trichloroethane	1,100,000(40,150)	200 <sup>(h)</sup>	-	-	1,900	100 <sup>(j)</sup>	Y
1,1,2-trichloroethane	82,500(25)	-	-	_	0.6	-	Y
Tetrachloroethene	50,000(4000)	-0.3	-	-	0.8	20 <sup>(j)</sup>	Y
Trichloroethene	275,000(100,000)	5 <sup>(h)</sup>	-	-	2.8	75(j)	Y
Toluene	133,750(28,250	-	-	15,000	-	<sup>75</sup> (j)	Y
Ethylbenzene	38,000(5,500)	-	-	24,000	-	-	Y
Phenol	3,950,000(76,250)	-	-	-	3,500	-	Y
PCB	75(25)	-	-	-	0.0006	-	Y

a All values in ug/L
b Safe Drinking Water Act Primary Maximum Contaminant Level
c Safe Drinking Water Act Secondary Maximum Contaminant Level
d Ambient Water Quality Criteria - Toxicity Protection
e Ambient Water Quality Criteria - 10-6 Cancer Risk
f Health advisory for protection of most sensitive population
g Organoleptic criteria
h Proposed MCL's
i MCL for trihalomethanes
c Chronic
k 10 Day

NOTE: Concentrations in ( ) are average projected release concentrations.

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Projected concentration of chloroform, trichloroethene, methylene chloride, tetrachloroethene, 1,1,1-trichoroethane, and toluene also would exceed standards and criteria.

#### Ingestion

Ingestion of groundwater could occur in both residential and occupational settings. Table 6-10 summarizes the risk assessment for the ingestion of the groundwater. In all settings, the excess lifetime cancer risk is greater than  $1 \times 10^{-6}$  with risk associated with projected concentrations in the shallow saturated zone exceeding  $1 \times 10^{-3}$ . Use of the shallow saturated zone and the shallow sand and gravel aquifer at the site could represent a potential public health risk without remedial action.

It is unlikely that the shallow saturated zone groundwater would be used as a water source due to the low hydraulic conductivity of this zone. The shallow confined aquifer would more likely be used. No new loadings into this zone are expected because of the upward gradient in this aquifer. It is possible that the concentration will decrease with time due to degradation. Because of that, the risk may be actually less.

Table 6-10
SUMMARY OF EXCESS LIFETIME CANCER RISK AND ACCEPTABLE DAILY INTAKE COMPARISONS
INGESTION OF GROUNDWATER AT THE ECC SITE

Contaminant Concentration Scenario	Setting	<u> Aquifer</u>	Major Chemical(s) of Concern	Total Excess Lifetime Cancer Risk	ADI _Exceeded ?
Current Values	Residential	Shallow Saturated Zone	Trichloroethene	2 x 10 <sup>-2</sup>	Trichloroethene
	Occupational	Shallow Saturated Zone	Trichloroethene	3 x 10 <sup>-3</sup>	Trichloroethene
Current Values	Residential	Shallow Sand and Gravel	1,1 Dichloroethene Tetrachloroethene Trichloroethene	7 x 10 <sup>-5</sup>	No
Current Values	Occupational	Shallow Sand and Gravel	1,1 Dichloroethene Tetrachloroethene Trichloroethene	1 x 10 <sup>-5</sup>	No
Projected Values (Maximum)	Residential	Shallow Saturated Zone	Methylene Chloride Tetrachloroethene Trichloroethene Chloroform PCB	4 x 10 <sup>-1</sup>	1,1,1-trichloroethane Toluene Ethylbenzene Phenol Trichloroethane Methylene chloride
Projected Values (Maximum)	Occupational	Shallow Saturated Zone	Methylene Chloride Tetrachloroethene Trichloroethene Chloroform PCB	8 x 10 <sup>-2</sup>	1,1,1-trichloroethane Toluene Ethylbenzene Phenol Trichloroethane Methylene chloride
Projected Values (Average)	Residential	Shallow Saturated Zone	Methylene Chloride Tetrachloroethene Trichloroethene Chloroform PCB	8 x 10 <sup>-3</sup>	1,1,1-trichloroethane Toluene Ethylbenzene Phenol Trichloroethane Methylene chloride
Projected Values (Average)	Occupational	Shallow Saturated Zone	Methylene Chloride Tetrachloroethene Trichloroethene Chloroform PCB	3 x 10 <sup>-3</sup>	1,1,1-trichloroethane Phenol Trichloroethane Methylene chloride

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#### Dermal Absorption

The dermal absorption of contaminants from groundwater would occur during bathing or showering. This would occur under the residential setting. Occupational showering and bathing would be very limited and is therefore not assessed.

A variety of factors can affect exposure from skin absorption including concentration, temperature, hydration of skin, duration and frequency of exposure. Skin absorption rates for most chemicals do not exist, and rates that do exist are for almost pure substances or high concentration aqueous solutions. The rates are often based on laboratory animal studies. While it is difficult to assess dermal absorption for many contaminants, it is possible to assess the absorption of volatile compounds (see Appendix D). The bathing risk estimation assumes that all of the compounds remain in the water phase and do not volatilize.

The risks are summarized in Table 6-11. The risk associated with bathing is roughly equal to the risk from ingestion and are greater than  $1 \times 10^{-6}$ . In both exposures, the volatile compounds are the chemicals of concern.

Under no action, bathing could represent a potential public health threat. However, by not accounting for volatilization, dermal absorption risks may be an overestimation.

# Table 6-11 SUMMARY OF EXCESS LIFETIME CANCER RISK DERMAL ABSORPTION OF GROUNDWATER AT THE ECC SITE

Contaminant Concentration Scenario	Aquifer	Major Chemical of Concern	Excess Lifetime Cancer Risk
Current Values	Shallow Sand and Gravel	1,1 Dichloroethene Trichloroethene	7 x 10 <sup>-5</sup>
Current Values	Shallow Saturated Zone	Trichloroethene	2 x 10 <sup>-2</sup>
Projected Values (Maximum)	Shallow Saturated Zone	Trichloroethene Methylene Chloride Tetrachloroethene Chloroform	4 x 10 <sup>-1</sup>
Projected Values (Average)	Shallow Saturated Zone	Trichloroethene Methylene Chloride Tetrachloroethene Chloroform	8 x 10 <sup>-2</sup>

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#### Vapor Inhalation

Vapors may be released from groundwater during use due to because of physical agitation of the groundwater or water temperatures raised above the volatilization point of the compound. This could occur in a variety of ways including bathing and cooking.

To model vapor release is difficult and information is lacking on human inhalation and retention efficiencies for individual chemical, therefore, it is not practical to estimate exposures and risk associated with this exposure route.

Only the qualitative statement that exposure may occur and increase risk can be made.

#### SURFACE WATER

The groundwater discharges to the unnamed ditch and Finley Creek. The surface water is a major release pathway for contaminants to leave the site. Fourteen compounds were found at the Finley Creek downstream sampling point (004) (see Tables 4-17 and 4-18). The volatile organic compounds are of greatest concern in terms of risk.

In addition to the measured concentrations in Finley Creek, it is possible to predict concentration in the surface water based on the projected concentration of contaminants in the

shallow saturated zone (from Chapter 5) and anticipated dilution with surface water. Dilutions are based on estimates of groundwater discharge to the unnamed ditch and Finley Creek, and USGS stream flow measurements for the unnamed ditch, Finley Creek, and Eagle Creek (see Table 6-4). Based on this information risks associated with exposures at the unnamed ditch, Finley Creek and Eagle Creek can be assessed.

The exposures that could occur at the surface waters would include direct exposure through wading via dermal absorption, and inhalation of volatile organics and indirect exposure by consumption of fish that have bioconcentrated contaminants from the surface water. Risks based on current concentrations in Finley Creek and projected concentrations in the unnamed ditch, Finley Creek, and Eagle Creek are assessed. These exposures are assessed detail in Appendix E and are summarized below.

#### Comparison to Standard

The current concentrations found in Finley Creek at SW004 and the projected concentration of contaminants in the unnamed ditch, Finley Creek and Eagle Creek are compared to the ambient water quality criteria for ingestion of aquatic organisms in Table 6-12. The concentration currently found at Finley do not exceed the criteria.

Table 6-12 COMPARISON OF SURFACE WATER CONCENTRATION TO AMBIENT WATER QUALITY CRITERIA FOR INGESTION OF AQUATIC ORGANISMS

	Current Concentration at SW004	Projected Concentration in Ditch		tration ey Creek	Projected Concentration in Eagle Creek ug/L	Ambient Water Quality Criteria-Ingestion of Aquatic Organism	
Compound	ug/L	ug/L	Maximum	Minimum	Maximum	ug/L	
1,1,1 Trichloroethane	120	67	33.5	1.7	0.8	1,030,000 <sup>a</sup>	
1,1 Dichloroethane	45	-	-	-	-	-	
Chloroethane	12	-	-	-	-	-	
1,2 Transdichloroethane	330	-	-	-	-	- h	
Tetrachloroethene	<5	7	3.5	0.18	0.09	8.85	
Trichloroethene	67	170	85	4.3	2.1	80.7	
Vinyl Chloride	10	-	-	-	-	525	
O-Xylene	<5	-	-	-	-	- h	
Methylene Chloride	-	170	85	4.3	2.1	15.7	
Toluene	-	50	25	1.3	0.6	424,000	
Aluminum	490	-	-	-	-	-	
Iron	1,410	-	-	-	-	-	
Manganese	130	-	-	-	-	-	
Cyanide	0.008	-	-	-	-	200 ີ	
1,1,2-trichloroethane	-	0.04	0.02	0.001	0.0005	41.8	
Phenol	-	130	65	3.3	1.6	769 <b>,</b> 000 b	
Chloroform	-	0.31	0.15	0.008	0.004	15.7	
Ethylbenzene	-	10	5.0	0.26	0.13	3,280	

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Based on toxicity.
Represents a 10 cancer risk level.

The projected concentrations do exceed the ambient water quality critera  $10^{-6}$  cancer risk for trichloroethene and methylene chloride in the unnamed ditch. The maximum projected concentration (i.e., lowest dilution) of methylene chloride and trichloroethene exceed the  $10^{-6}$  level in Finley Creek.

#### Dermal Absorption

Residents and visitors could be exposed to volatile chemicals in the surface water by wading in the unnamed ditch, Finley Creek and Eagle Creek during the warmer months of the year. Assumptions concerning wading appear in Appendix D. The actual population currently at risk is unknown but expected to be small. The area is growing and the population exposed could increase. The risks are summarized in Table 6-13. Wading in these waterways does not exceed  $1 \times 10^{-6}$  excess lifetime cancer risk.

#### Ingestion Via Fish Consumption

Fish have been observed in Finley and Eagle Creek. Human exposure to contaminants could occur from consumption of fish that are caught if the fish have bioconcentrated surface water contamiants. There were no fish samples taken, therefore, literature values for bioconcentration factors are used.

## Table 6-13 SUMMARY OF EXCESS LIFETIME CANCER RISK FROM WADING - ECC

#### CURRENT CONDITIONS

Location	Risk
Finley Creek	$5 \times 10^{-7}$
PREDICTED CONDITIONS <sup>a</sup>	
Location	Risk
Unnamed Ditch	$6 \times 10^{-7} b$
Finley Creek	$4 \times 10^{-7} \text{ c}$

 $1 \times 10^{-8} d$ 

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Eagle Creek

aBased upon the projected contaminant concentrations released to the groundwater from the soil.

Assume 1:600 groundwater to ditch water dilution.

Assume 1:2 ditch to Finley Creek dilution.

Assume 1:41 Finley Creek to Eagle Creek dilute.

The current concentration measured in Finley Creek and as the projected concentrations for the unnamed ditch, Finley Creek, and Eagle Creek are assessed. The projected discharge of PCB to the surface water is not included in the assessment because the time frame for the migration of PCB's from soil to surface water via groundwater discharge would be orders-of-magnitude greater than the other compounds. The results are summarized in Table 6-14.

The excess lifetime cancer risk from fish ingestion under the current concentrations in Finley Creek is  $1 \times 10^{-6}$ . The projected values for the unnamed ditch and Finley Creek (under the least dilution) are slightly greater than  $1 \times 10^{-6}$ .

This risk estimation relies on a number of assumptions (see Appendix E and Table 6-4) and projected values such that the risks presented represent a conservative upper bound. It is unlikely that a sufficient number of fish are residing in the unnamed ditch to make the analysis realistic. It is also unlikely that both fish and fishermen would be restricted to one stream segment. The approach that is taken, is taken for simplicity sake and it's limitations are recognized.

Table 6-14
SUMMARY OF EXCESS LIFETIME CANCER RISK
FROM CONSUMPTION OF FISH IN THE WATERWAYS AT THE ECC SITE

Location	Scenario	Risk
Finley Creek	Actual Concentration	1 x 10 <sup>-6</sup>
Unnamed Ditch	Projected Concentration	$3 \times 10^{-6}$
Finley Creek	Projected Concentration (Least dilution)	$2 \times 10^{-6}$
Finley Creek	Projected Concentration (Greatest dilution)	$1 \times 10^{-7}$
Eagle Creek	Projected Concentration (Least dilution)	$4 \times 10^{-8}$

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#### ENVIRONMENTAL ASSESSMENT

#### INTRODUCTION

This environmental assessment describes the current site situation and the environmental conditions anticipated if no remedial action is taken. This assessment identifies habitats that are or could become contaminated, the types of impacts that are likely and assesses the general significance of the impacts.

#### Population at Risk

The population at risk would be the terrestrial and aquatic animal species and associated plant communities that reside on or include the ECC site and adjacent areas as part of their range. This would include species that permanently reside in the area as well as transient species. The population at risk and their route of exposure include:

- o Aquatic organisms, through contamination of surface waters from runoff or discharges into them.
- Local vegetation through contact with contaminated sediment or dust.

o Local fish, wildlife, and domestic animals,
through contact with or ingestion of contaminated
vegetation, soil, sediment, or surface water.

The area is former agricultural land with second growth plant communities in the fields and dense plant growth along the waterways. The ECC site drains into the riverine type wetlands that are comprised of the unnamed ditch, Finley Creek and Eagle Creek. There are no known designated critical habitats for threatened or endangered species that are impacted by the ECC site. There are no known endangered species that inhabit the area around the ECC site.

Several of the compounds, trichloroethene and tetrachloroethene are known to bioconcentrate. Food chain affects could occur if fish are eaten by terrestrial organisms.

#### SOIL AND SEDIMENT

Some of the organic contaminants found in soil and sedimebnt bioaccumulate and tend to stay in the fatty tissue of animals once ingested. Eight of the inorganics found in the soil (arsenic, cadmium, chromium, copper, cyanide, nickel, lead, and mercury) and three of the inorganics found in the sediment (cyanide, mercury, and lead) tend to adsorb on clay and organic particles in the soil or sediment which

ultimately may be deposited on plants as dust. Animals may also inadvertently ingest contaminated soil or exposed sediment while grooming and feeding. Some of the compounds may be taken up by plants and ultimately eaten by animals both of which may or may not be adversely affected.

#### SURFACE WATER

The discharge of contaminanted groundwater would have the greatest potential impact on the aquatic environments. To a lesser extent surface runoff would also affect the aquatic environments. The Depauw University study on trophic composition of the fish population suggests an impact on the aquatic population in Finley Creek (see Chapter 3). This impact can not be conclusively associated with the ECC site, however. The State of Indiana's mussel bioaccumulation study was inconclusive (see Chapter 3).

Table 6-15 compares concentrations found at sampling point 004 and projected concentrations in the unnamed ditch, Finley Creek and Eagle Creek to ambient water quality criteria and 96 hour  ${\rm LC}_{50}$  values. Concentrations do not exceed either  ${\rm LC}_{50}$  values or water quality criteria for protection of aquifer life under any of the conditions assessed .

Table 6-15 COMPARISON TO AMBIENT WATER QUALITY CRITERIA AND 96 HR LC 50

Compound	Finley 004 Concentration ug/L	Projected Unnamed Ditch Concentration ug/L	Maximum Projected Finley Creek Concentration ug/L	Maximum Projected Eagle Creek Concentration ug/L	•	pc <sup>C</sup> Protection  /L   Chronic	96 hr LC ug7L
1,1,1 Trichloroethane	120	67	34	0.85	18,000		52,800 a
1,1 Dichloroethane	45				••		550,000 <sup>D</sup>
Trans 1,2 Dichloroethene					16,000		
Methylene Chloride	<5	170	85	2.1			193,000
Tetrachloroethene	<5	7	3.5	0.09	5,280	840	18,400 <sup>a</sup>
Trichloroethene	620	170	85	2.1	45,000		40,200ª
Vinyl Chloride	10						
Xylene	5						42,000 <sup>a</sup>
Toluene		50	25	0.62	17,500		34,000 <sup>a</sup>
Phenol		130	65	1.6	10,200 <sup>c</sup>	2,560 <sup>C</sup>	5,700 <sup>D</sup>
Ethylbenzene		10	5	0.12	32,000		42,300 <sup>a</sup>

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a For flathead minnow b For bluegill C Ambient Water Quality Criteria listing of lowest adverse effects on aquatic life d Lethal concentration 50% over 96 hour period

#### SUMMARY

The major public health and environmental risks from the ECC site derived in this endangerment assessment are outlined in Table 6-16. Each risk is listed by pathway and the likelihood of the risk is assessed. The major risks come from the contaminated soil via direct contact and release of soil contaminants to the groundwater and subsequent use of groundwater for bathing and drinking water source. The current population at risk is limited and while the area is projected to grow the impact of the ECC site appears to be localized.

In conclusion, the site does pose a potential threat to the public health, welfare, and environment, and a feasibility study of remedial action to cost-effectively mitigate the site hazards should be performed.

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### Table 6-16 (Page 1 of 4) SUMMARY OF MAJOR RISK FROM ENDANGERMENT ASSESSMENT RISK/EFFECTS

Pathway	Location	Setting	Excess Lifetime Cancer Risk	Acceptable Daily Intake (ADI)	Compounds of Concern	Comment	Probability
Public Health Evaluat	ion						
Soil - Direct contact via ingestion	South Pad - Intermediate Depth	Residential	4 x 10 <sup>-5</sup> to 8 x 10 <sup>-6</sup>	•	Trichloroethene Tetrachloroethene	Based on maximum to average concentration	Requires development of site - limited area of exposure.
Soil - Direct contact via ingestion	Northern Test Pit Area - Shallow Depth	Residential	4 x 10 <sup>-3</sup> to 3 x 10 <sup>-4</sup>	-	PCB's Trichloroethene Tetrachloroethene	Based on maximum to average concentration	Requires development of site - limited area of exposure.
Soil - Direct contact via ingestion	Northern Test Pit Area - Shallow Depth	Residential	-	ADI's exceeded at 1 gram/day ingestion rate	Xylene Lead	Based on maximum concentrations	Requires development of site - limited area of exposure.
Soil - Direct contact via ingestion	Northern Test Pit Area - Intermediate Depth	Residential	8 x 10 <sup>-4</sup> to 2 x 10 <sup>-5</sup>	-	PCB's Trichloroethene Tetrachloroethene	Based on maximum to average concentration	Requires development of site - limited area of exposure.
Soil - Direct contact via ingestion	Northern Test Pit Area - Intermediate Depth	Residential	-	ADI's exceeded at 10 grams/day ADI exceeded at 1 gram/day	Cadmium Lead Lead	Based on maximum concentrations	Requires development of site - limited area of exposure.
Sediment - Direct Contact via ingestion	Finley Creek downstream from ECC at high- way 421	Residential	-	ADI exceeded at 10 gram/day	Lead	Based on maximum concentration	Requires exposure of or direct contact with sediment. Season- ally limited. Contamination cannot be directly associated with the ECC site.
Groundwater - via ingestion	Onsite - Shallow Saturated Zone	Residential - current contam- inant levels	2 x 10 <sup>-2</sup> to 3 x 10 <sup>-3</sup>	ADI exceeded at 10 gram/day	Trichloroethene	Based on one sampling point	No current exposures. Requires development of site. Potential future exposed population limited by size of area and low permeability of water bearing soil. Contaminant levels may increase with time.

							•
Pathway  Groundwater - via ingestion	Location  Onsite - Shallow Sand and Gravel Aquifer	Residential Occupational current contam- inant levels	Excess Lifetime Cancer Risk  7 x 10 <sup>-5</sup> to 1 x 10	Acceptable Daily Intake (ADI)	Compounds of Concern  1,1-Dichloroethene Tetrachloroethene Trichloroethene	Comment  Based on maximum concentrations	Probability  No current exposures based on residential well data. Limited potential future exposed population. Upward gradient should limit new contamination - concentration and risk should decline with time.
Groundwater - via ingestion	Onsite/Offsite Shallow Saturated Zone	Residential Occupational Project Maximum Values	4 x 10 <sup>-1</sup> 8 x 10 <sup>-2</sup>	ADI exceeded ADI exceeded	Methylene Chloride Tetrachloroethene Trichloroethene Chloroform PCB	Based on projected release from soil, no degradation and maximum concentra- tion	Requires development of site surrounding area. Upper bound value based on highest soil concentrations. Actual population using groundwater would be limited by size of area and low permeability of water bearing soil.
Groundwater - via ingestion	Onsite/Offsite Shallow Saturated Zone	Residential Occupational Project Average Values	8 x 10 <sup>-3</sup> 3 x 10 <sup>-3</sup>	ADI exceeded ADI exceeded	Methylene Chloride Tetrachloroethene Trichloroethene Chloroform PCB	Based on projected release from soil, no degradation and average concentra- tion	Requires development of site surrounding area. Upper bound value based on highest soil concentrations.
Groundwater - via dermal absorption (bathing)	Onsite - Water table aquifer	Residential Current contam- inant levels	2 x 10 <sup>-2</sup>	-	Trichloroethene	Based on one sampling point. Assumes no volatilization.	No current exposures. Requires development of site. Potential future exposed population limited by size of area and low permeability of water bearing soil. Contaminant levels may increase with time.
Groundwater - via dermal absorption (bathing)	Onsite - Shallow confined aquifer	Residential - Current contam- inant levels	7 x 10 <sup>-7</sup>	-	1,1-Dichloroethene Trichloroethene	Based on maximum concentrations.  Assumes no volatilization.	No current exposures based on residential well data. Limited potential future exposed population. Upwared gradient should limit new contamination concentration and risk should decline with time.
Groundwater - via dermal absorption (bathing)	Onsite/offsite water table aquifer	Residential Projected Maximum values	4 x 10 <sup>-1</sup>	-	Methylene Chloride Tetrachloroethene Trichloroethene Chloroform	Based on projected release from soil, no degradation, no volatilization and maximum con- centration	Requires development of site/ surrounding area. Upper bound value based on highest soil con- centrations. Actual population using groundwater would be limited by size of area and low permeability of water bearing soil.

Pathway	Location	Setting	Excess Lifetime Cancer Risk	Acceptable Daily Intake (ADI)	Compounds of Concern	Comment	Probability
Groundwater - via dermal absorption (bathing)	Onsite/offsite water table aquifer	Residential Projected Average values	8 x 10 <sup>-2</sup>	-	Methylene Chloride Tetrachloroethene Trichloroethene Chloroform	Based on projected release from soil, no degradation, no volatilization	Requires development of site/ surrounding area. Upper bound value based on highest soil concentrations.
Groundwater dis- charge to surface water - dermal absorption from wading	Finley Creek	Actual Concentrations	5 x 10 <sup>-7</sup>	-	Trichloroethene	Based on one sampling point	Assumes concentrations remain constant. Cannot be definitly associated with ECC. Limited potential of exposed population.
Groundwater dis- charge to surface water - dermai absorption from wading	Unnamed Ditch Finley Creek Eagle Creek	Projected Concentrations	6 x 10 <sup>-7</sup> 6 x 10 <sup>-7</sup> 6 x 10 <sup>-8</sup>	-	Trichloroethene Tetrachloroethene Methylene Chloride	Based on projected concentrations over a range of dilutions. Assumes no volatilization.	Upper bound range of risk based on average release from soil. Limited potentially exposed population.
Groundwater dis- charge to surface water - fish bio- concentration of contaminants - human ingestion of fish	Finley Creek	Actual Concentrations	1 x 10 <sup>-6</sup>	-	Trichloroethene Tetrachloroethene	Based on one sampling point and literature values for BCF	Assumes: exclusive and active fishing in Finley Creek; fish reside exclusive in Finley Creek; sufficient sport fish population. Currently exposed population unknown but estimated to be small. Some comments for future.
Groundwater dis- charge to surface water - fish bio- concentration of contaminants - human ingestion of fish	Unnamed Ditch	Projected Concentrations	3 x 10 <sup>-6</sup>	-	Trichloroethene Tetrachloroethene Methylene Chloride Chloroform	Based on projected concentrations over a range of dilutions. Uses average soil concentration as a basis. Assumes no volatilization. Based on literature values for BCF.	Values are upper bound range. Exposed population unknown but estimated to be small. Assumes: exclusive and active fishing in creek; fish reside exclusively in creek; sufficient sport fish population. Volatilization should reduce concentration.

Pathway	Location	Setting	Excess Lifetime Cancer Risk	Acceptable Daily Intake (ADI)	Compounds of Concern	Comment	Probability
Groundwater dis- charge to surface water - fish bio- concentration of contaminants - human ingestion of fish	Finley Creek	Projected Concentrations	2 x 10 <sup>-6</sup> to 1 x 10 <sup>-7</sup>	-	Trichloroethene Tetrachloroethene Methylene Chloride Chloroform	Based on projected concentrations over a range of dilutions. Uses average soil concentration as a basis. Assumes no volatilization. Based on literature values for BCF.	Values are upper bound range. Exposed population unknown but estimated to be small. Assumes: exclusive and active fishing in creek; fish reside exclusively in creek; sufficient sport fish population. Volatilization should reduce concentration.
Groundwater dis- charge to surface water - fish blo- concentration of contaminants - human ingestion of fish	Eagle Creek	Projected Concentrations	4 x 10 <sup>-8</sup>	-	Trichloroethene Tetrachloroethene Methylene Chloride Chloroform	Based on projected concentrations the least of dilutions. Uses average soil concentration as a basis. Assumes no volatilization. Based on literature values for BCF.	Values are upper bound range. Exposed population unknown but estimated to be small. Assumes: exclusive and active fishing in creek; fish reside exclusively in creek; sufficient sport fish population. Volatilization should reduce concentration.

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